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(71) Applicant: KYOWA HAKKO KOGYO CO., LTD. Chiyoda-ku, Tokyo 100 (JP)

(72) Inventors:

· KOIKE, Masamichi Machida-shi, Tokyo 194 (JP)

 FURUYA, Akiko Machida-shi, Tokyo 194 (JP) NAKAMURA, Kazuyasu Machida-shi, Tokyo 194 (JP)

 IIDA, Akihiro Machida-shi, Tokyo 194 (JP)

· ANAZAWA, Hideharu Tokyo 178 (JP)

 HANAI, Nobuo Sagamihara-shi, Kanagawa 229 (JP)

• TAKATSU, Kiyoshi Tokyo 113 (JP)

(74) Representative: **VOSSIUS & PARTNER** Postfach 86 07 67 81634 München (DE)

ANTIBODY AGAINTS ALPHA-CHAIN OF HUMAN INTERLEUKIN 5 RECEPTOR (54)

The present invention provides monoclonal antibodies and humanized antibodies which react specifically with a human interleukin-5 receptor α chain. The invention also provides hybridomas and transformants which produce the antibodies, the monoclonal antibodies and humanized antibodies, a method for detecting an interleukin-5 receptor α chain immunologically by means of these antibodies, as well as a method for diagnosing and treating diseases such as chronic bronchial asthma by means of the monoclonal antibodies and humanized antibodies. The present invention is useful for diagnosis or treatment of diseases such as chronic bronchial asthma.

EP 0 811 691 A1

Description

Technical Field

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The present invention relates to monoclonal antibodies and humanized antibodies which bind specifically to a human interleukin-5 receptor α chain and which are therefore useful for diagnosis or treatment of diseases such as chronic bronchial asthma. The invention also relates to hybridomas and transformants which produce the antibodies, a method for detecting an interleukin-5 receptor α chain immunologically by means of the monoclonal antibodies and humanized antibodies, as well as a method for diagnosing and treating diseases such as chronic bronchial asthma by means of the monoclonal antibodies and humanized antibodies.

Background Art

Interleukin-5 (hereinafter referred to as "IL-5") is a kind of lymphokine which is secreted by T cells, mast cells and other cells. Murine IL-5 is known to act as a differentiation and growth factor for B cells and eosinophils. Human IL-5 is known to act mainly as a differentiation and growth factor for eosinophils (Advances in Immunology, 57, 145 (1994); Blood, 79, 3101 (1992)). IL-5 exhibits its action through a specific receptor (IL-5 receptor) which is expressed on the surface of a cell such as eosinophil. It has been shown that human and murine IL-5 receptors (hereinafter referred to as "IL-5Rs") are both composed of two different kinds of proteins, an α chain (hereinafter referred to as "IL-5R α ") and a β chain (hereinafter referred to as "IL-5R β "). In addition, it is known that the binding of IL-5 to IL-5R is via IL-5R α and that IL-5R β alone can not bind to IL-5 (EMBO J., 9, 4367 (1990); ibid., 10, 2833 (1991); J. Exp. Med., 177, 1523 (1993); ibid., 175, 341 (1992); Cell, 175, 175 (1991), Proc. Natl. Acad. Sci., 175,

Eosinophils are known to increase in allergic diseases represented by chronic bronchial asthma. Significant infiltration of eosinophils is observed in airways of a patient with chronic bronchial asthma. Eosinophil contains a cytotoxic granular proteins whose deposit is observed in airway tissues of a patient with chronic bronchial asthma or at lesion sites of a patient with atopic dermatitis. These facts suggest that eosinophil plays an important role in the pathogenesis of allergic disorders such as chronic bronchial asthma, atopic dermatitis and the like (Adv. Immunol., 39, 177 (1986); Immunol. Today, 13, 501 (1992)). Hence, studying the kinetics of eosinophils is useful for clinical diagnosis. On the other hand, human IL-5 acts specifically on eosinophils, so IL-5R is believed to be expressed specifically in eosinophils and can therefore be used as a marker specific to human eosinophils. Furthermore, IL-5 β is a receptor for cytokines such as IL-3, GM-CSF and others, so IL-5R α is believed to be a marker specific to eosinophils. Hence, eosinophils can be detected specifically by immunocyte staining using an anti-human IL-5R α chain antibody (hereinafter referred to as "anti-hIL-5R α antibody"). However, no anti-hIL-5R α antibody is presently known that is capable of specific detection of eosinophils.

Significant eosinophilia was observed in IL-5 transgenic mice (J. Exp. Med., <u>172</u>, 1425 (1990); ibid. <u>173</u>, 429 (1991); Int. Immunol., <u>2</u>, 965 (1990)). Eosinophil infiltration in tissues was suppressed by the administration of an anti-IL-5 antibody in animal models of asthma (Am. Rev. Resir. <u>147</u>, 548 (1993); ibid., <u>148</u>, 1623 (1993)). These phenomena indicate that IL-5 actually plays an important role in eosinophilia and the infiltration of eosinophils in vivo. It is also reported that IL-5 is expressed in airway mucosal tissues of a human patient with chronic bronchial asthma and at lesion sites of a patient with atopic dermatitis (J. Clin. Invest., <u>87</u>, 1541 (1991); J. Exp. Med., <u>173</u>, 775 (1991)). Further investigations demonstrate that IL-5 exhibits in vitro viability-enhancing action on human eosinophils (J. Immunol., <u>143</u>, 2311 (1989)) and that IL-5 is an eosinophil-selective activator (J. Exp. Med., <u>167</u>, 219 (1988)).

Hence, antibodies that bind to IL-5R and which can inhibit the biological activity of IL-5 are expected to inhibit the activity of eosinophil, thus being useful in the treatment of allergic diseases such as chronic bronchial asthma. Antimouse IL-5R α antibodies which can inhibit the biological activity of IL-5 were produced by using as an antigen those IL-5-dependent cells which express a large number of murine IL-5R on their surfaces (Kokai (Japanese published unexamined patent application) No. 108497/91; Int. Immunol., 2, 181 (1990)). However, in the case of humans, no cells are known which express a large number of IL-5R and the expression of IL-5R is reported to be very low in eosinophils (Cell. Immunol., 133, 484 (1991)). Hence, anti-human IL-5R α antibodies having comparable functions to anti-mouse IL-5R α antibodies are difficult to produce by methods similar to those for producing the latter. An antibody designated as " α 16" is disclosed as an antibody against human IL-5R α in EMBO J., 14, 3395 (1995) but this antibody does not have any neutralization activity for IL-5R α .

Human IL-5R α gene was obtained by preparing a cDNA library from human eosinophil (J. Exp. Med., <u>175</u>, 341 (1992)) or a human promyelocytic cell HL-60 (Cell, <u>66</u>, 1175 (1991); Kokai No. 78772/94) and screening the library using as a probe an oligo DNA which had been synthesized on the basis of cDNA of murine IL-5R α or a partial amino acid sequence of murine IL-5R α (Kokai No. 54690/94, EMBO J., <u>9</u>, 4367 (1990)). The transfer of the cDNA into a host

cell resulted in the creation of a cell having hIL-5R α expressed on its surface but the expression level of hIL-5R in this cell was very low ($\leq 10^4$ molecules) (J. Exp. Med., <u>177</u>, 1523 (1993)). Hence, if one attempts to produce anti-hIL-5R α antibodies by using this cell as an immunogen, he will find that the relative amount of hIL-5R α is very small, compared with those of proteins from a host cell and that the absolute protein amount of hIL-5R α is also very small. In addition, approximately 80% homology at an amino acid level is observed between murine IL-5R α and human IL-5R α and murine IL-5 can bind to human IL-5R with high affinity (J. Exp. Med., <u>175</u>, 341 (1992)). These facts suggest that human IL-5R α has a lower immunogenicity for mice or rats which are commonly used as animals to be immunized. In fact, almost all of our attempts to prepare anti-hIL-5R α antibodies using hIL-5R α - expressing cells as an immunogen resulted in a failure.

In the cloning of IL-5R cDNA from a cDNA library of human eosinophil, cDNA encoding soluble human IL-5R α (hereinafter referred to as "shIL-5R α) has been obtained which corresponds to the N-terminal amino acid sequence (1-313) of IL-5R α which is defective in the transmembrane region and onwards (J, Exp. Med.,175, 341 (1992)). When shIL-5R α is used as an immunogen to produce an anti-hIL-5R α antibody, the shIL-5R α should have the same three dimensional conformation as that of IL-5R α expressed on the cell surface and it should be one secreted and produced by a eukaryotic host cell in order to obtain an anti-hIL-5R α antibody which can inhibit the biological activity of IL-5. In addition, it has been found that the production efficiency of a protein varies significantly depending on the signal peptide (Protein, Nucleic Acid and Enzyme,35, 2584 (1990)), so it is necessary to select an appropriate signal peptide for secretion and production of the protein.

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As mentioned above, it has been found that mRNA which is believed to encode only shIL-5R α is expressed in eosinophils. It has been confirmed that murine IL-5R is expressed not only in eosinophils but also in B cells and that mRNA which is believed to encode only an extracellular region of IL-5R α (hereinafter referred to as "smIL-5R α) is expressed in those cells as well as in the case of humans. In addition, it has been reported that smIL-5R α was detected in blood of mice transplanted with IL-5R expressing murine chronic B cell leukemia cell line (BCL1) or model mice of human autoimmune diseases (J. Immunol. Method, 167, 289 (1994)). These suggest the possibility that the increase in the number of IL-5R expressing cells and their activation may be reflected in the amount of smIL-5R α secreted in blood. Human IL-5R is believed to be expressed in eosinophils in a limited amount and the increase in the number of eosinophils and their activation may be potentially reflected in the amount of shIL-5R α in blood. Hence, the quantitative determination of shIL-5R α is expected to be useful in clinical diagnosis.

Any isolated monoclonal antibody which binds specifically to human IL-5R α is believed to be useful in the diagnosis and treatment of allergic diseases. However, it should be noted that if a non-human animal-derived monoclonal antibody is administered to a human, it is generally recognized as a foreign matter such that an antibody against the non-human animal-derived monoclonal antibody is produced in the human body, a reaction with the administered non-human animal-derived monoclonal antibody occurs to cause a side effect (J. Clin. Oncol., $\underline{2}$, 881 (1984); Blood, $\underline{65}$, 1349 (1985); J. Natl. Cancer Inst., $\underline{80}$, 932 (1988); Proc. Natl. Acad. Sci., $\underline{82}$, 1242 (1985)), premature clearance of the non-human animal-derived monoclonal antibody occurs (J. Nucl. Med., $\underline{26}$, 1011 (1985); Blood, $\underline{65}$, 1349 (1985); J. Natl. Cancer Inst., $\underline{80}$, 937 (1988)), or therapeutic effect of the monoclonal antibody is reduced (J. Immunol., $\underline{135}$, 1530 (1985); Cancer Res., $\underline{46}$, 6489 (1986)).

In order to solve these problems, attempts have been made to convert non-human animal-derived monoclonal antibodies to human chimeric antibodies or human CDR-grafted antibodies (reconstituted human antibodies) by gene recombinant techniques. A human chimeric antibody is an antibody of which the variable region (hereinafter referred to as "V region") is derived from a non-human animal antibody and the constant region (hereinafter referred to as "C region") is derived from a human antibody (Proc. Natl. Acad. Sci., <u>81</u>, 6851 (1984)). It has been reported that when a human chimeric antibody is administered to a human, antibodies are hardly produced against the non-human animal-derived monoclonal antibody and a half-life in blood is increased by a factor of 6 (Proc. Natl. Acad. Sci., <u>86</u>, 4220 (1989)). A human CDR-grafted antibody is a human antibody of which the CDR (complementarity determining region) is replaced with the CDR of a nonhuman animal-derived antibody (Nature, <u>321</u>, 522 (1986)). It has been reported with experiments on monkeys that a human CDR-grafted antibody has a lower immunogenicity, with the half-life in blood being increased by a factor of 4-5 compared with a mouse antibody (J. Immunol., <u>147</u>, 1352 (1991)). However, there is no report about a humanized antibody against hIL-5R α.

When a humanized antibody which binds specifically to human IL-5R α is administered to a human, it is expected to cause no production of an antibody against a non-human animal-derived monoclonal antibody, thereby reducing the side effect and prolonging the half-life in blood, which eventually leads to a high therapeutic effect against allergic diseases such as chronic bronchial asthma, atopic dermatitis and the like.

As a result or the recent progresses in protein and genetic engineerings, smaller antibody molecules such as single chain antibodies (Science, <u>242</u>, 423 (1988)) and disulfide stabilized antibodies (Molecular Immunology, <u>32</u>, 249 (1995)) are being prepared. Since single chain antibodies and disulfide stabilized antibodies have smaller molecular weights than monoclonal antibodies and humanized antibodies, they are effective in transition into tissues and clearance from blood and their application to the imaging technology and the preparation of complexes with toxins are being underway to provide some promise in therapeutic efficacy (Cancer Research, <u>55</u>, 318 (1995)). If a single chain antibody or a

disulfide stabilized antibody which binds specifically to a human IL-5R α chain is produced, high diagnostic and therapeutic effects against allergic diseases and the like are anticipated. However, there is no report about a single chain antibody and a disulfide stabilized antibody against a human IL-5R α chain.

5 <u>Disclosure of Invention</u>

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The inventors found that antibodies to a hIL-5R α chain which recognizes an epitope at 1-313 positions of the N-terminal amino acid sequence of the human IL-5R α chain which corresponds to an extracellular region defective in the transmembrane region and onwards react specifically with a human interleukin-5 receptor α chain upon immunocyte staining and inhibit the biological activity of interleukin-5. These antibodies can be used to diagnose and treat the aforementioned allergic diseases.

Hence, the present invention provides antibodies which react specifically with a human IL-5R α chain. The antibodies of the present invention include monoclonal antibodies, humanized antibodies, single chain antibodies, disulfide stabilized antibodies and the like. The antibodies or the present invention may be of any kinds, provided that they react specifically with a hIL-5R α chain. Those produced by the method explained below are preferred. Briefly, hIL-5R α protein is prepared as an antigen and applied to immunize animals such as mice, rats, hamsters, rabbits and the like used to prepare hybridomas, thereby inducing to plasma cells having an antigen specificity. The plasma cells are fused with myeloma cells to prepare hybridomas which can produce monoclonal antibodies, and the hybridomas are cultured to obtain the desired anti-IL-5R α monoclonal antibodies. Any monoclonal antibodies can be used so long as it recognizes an epitope at 1-313 positions from the N-terminal amino acid of a human IL-5R α chain and reacts specifically with the human IL-5R α chain upon immunocyte staining. Alternatively, any monoclonal antibodies can be used so long as it recognizes an epitope at 1-313 positions from the N-terminal amino acid of the human IL-5R α chain and inhibits the biological activity of human IL-5. The former monoclonal antibodies are exemplified by monoclonal antibody KM1257 produced by hybridoma KM1257 (FERM BP-5133). The latter monoclonal antibodies are exemplified by KM1259 produced by hybridoma KM1259 (FERM BP-5134) and KM1486 produced by hybridoma KM1486 (FERM BP-5651).

The monoclonal antibodies of the present invention react immunologically with a human IL-5R α chain, a cell having a human IL-5R α chain expressed on the surface, human eosinophil and the like. The monoclonal antibodies of the present invention react immunologically with a soluble human IL-5R α chain. Hence, the present invention also provides a method for immunologically detecting and determining a human IL-5R α chain, a cell having a human IL-5R α chain expressed on the surface, human eosinophil and a soluble human IL-5R α chain. The results of the detection and determination can be used in the diagnosis and treatment of allergic diseases such as chronic bronchial asthma, atopic dermatitis and the like.

The present invention also provides humanized antibodies that have lesser side effects with a prolonged half-life than the monoclonal antibodies and which inhibit the biological activity of IL-5 in a more desired way as therapeutics. The term "humanized antibody" of the present invention is the general term for human chimeric antibodies and human CDR-grafted antibodies.

The term "human chimeric antibody" means an antibody consisting of a variable region in a heavy chain (hereinafter referred to as "VH") and a variable region in a light chain (hereinafter referred to as "VH") of a non-human animal antibody, as well as a constant region in a heavy chain (hereinafter referred to as "CH") and a constant region in a light chain (hereinafter referred to as "CH") of a human antibody. The term "human CDR-grafted antibody" means an antibody in which CDR sequences of VH and VL of a human antibody are replaced with CDR sequences of VH and VL of a non-human animal antibody, respectively. An anti-hIL-5R α chain human chimeric antibody which inhibits the biological activity of IL-5 can be expressed and produced by a process comprising the steps of obtaining cDNAs encoding VH and VL from a hybridoma producing an antibody which can inhibit the biological activity of IL-5, inserting the respective cDNAs into a vector for expression in animal cells which contains a gene encoding human antibody CH and human antibody CL to thereby construct a human chimeric antibody expression vector and transfecting the expression vector into an animal cell. The human chimeric antibody and human CDR-grafted antibody of the present invention may be in any immunoglobulin (Ig) classes and are preferably in a class of IgG. In addition, any C region of IgG subclasses of immunoglobulin such as IgG1, IgG2, IgG3 and IgG4 can be used.

Examples of the human chimeric antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 24, CH is human antibody IgG1, VL contains the amino acid sequence of SEQ ID NO: 25, and CL is human antibody κ. A specific example is an antibody designated as "KM1399". A specific example of the human chimeric antibody of which the CH is human antibody IgG4 is an antibody designated as "KM7399". KM1399 can be produced, for example, by transformant KM1399 (FERM BP-5650). KM7399 can be produced, for example, by transformant KM7399 (FERM BP-5649).

The anti-hIL-5R α chain human CDR-grafted antibody which inhibits the biological activity of IL-5 can be expressed and produced by a process comprising the steps of constructing cDNAs encoding a V region in which CDR sequences of VH and VL of any human antibody are replaced with CDR sequences of VH and VL, respectively, of a non-human animal antibody which can inhibit the biological activity of IL-5, inserting the respective cDNAs into a vector for expres-

sion in animal cells which contains a gene encoding human antibody CH and human antibody CL to thereby construct a human CDR-grafted antibody expression vector, and transfecting the expression vector into an animal cell. Examples of the human CDR-grafted antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 72, CH is human antibody IgG1, VL contains the amino acid sequence of SEQ ID NO: 63, and CL is human antibody κ . A specific example is an antibody designated as "KM8399". A specific example of the human CDR-grafted antibody of which the CH is human antibody IgG4 is an antibody designated as "KM9399". KM8399 can be produced, for example, by transformant KM8399 (FERM BP-5648). KM9399 can be produced, for example, by transformant KM9399 (FERM BP-5647).

The humanized antibody of the present invention reacts immunologically with a human IL-5R α chain, a cell having a human IL-5R α chain expressed on the surface, human eosinophil and the like. Hence, the humanized antibody of the present invention can be used in the diagnosis and treatment of allergic diseases such as chronic bronchial asthma, atopic dermatitis and the like.

In addition, the present invention provides single chain antibodies (single chain Fv; hereinafter referred to as "scFv") and disulfide stabilized antibodies (disulfide stabilized Fv; hereinafter referred to as "dsFv") which exhibit an ability to bind to a human IL-5R α chain.

The term "single chain antibody (scFv)" means a polypeptide represented by formula VH-L-VL or VL-L-VH in which a single chain of VH and a single chain of VL are linked by an appropriate peptide linker (hereinafter referred to as "L"). Any anti-human IL-5R α chain monoclonal antibodies or human CDR-grafted antibodies can be used as VH and VL in the scFv of the present invention.

The term "disulfide stabilized antibody (dsFv)" means an antibody prepared by binding through a disulfide bond two polypeptides in which each one of the amino acid residues in VH and VL is replaced with cysteine residues. The amino acid residues to be replaced with cysteine residues can be selected on the basis of a presumed steric structure of an antibody in accordance with the method described by Reiter et al. (Protein Engineering, \underline{Z} , 697 (1994)). Either a mouse anti-human IL-5R α chain monoclonal antibodies or a human CDR-grafted antibodies can be used as VH or VL in the disulfide stabilized antibody of the present invention.

The single chain antibody which has an ability to bind to a human IL-5R α chain can be expressed and produced by a process comprising the steps of obtaining cDNA encoding VH and VL from a hybridoma which produces an antibody reactive with the human IL-5R α chain, constructing a single chain antibody expression vector, and transfecting the expression vector into an <u>E. coli</u>, yeast or animal cell. Examples of the monoclonal antibody-derived single chain antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 24 and VL contains the amino acid sequence of SEQ ID NO: 25. Examples of the human CDR-grafted antibody-derived single chain antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 72 and VL contains the amino acid sequence of SEQ ID NO: 63.

The disulfide stabilized antibody which has an ability to bind to a human IL-5R α chain can be expressed and produced by a process comprising the steps of obtaining cDNA encoding VH and VL from a hybridoma which produces an antibody reactive with the human IL-5R α chain, inserting the cDNA into an appropriate expression vector, and transfecting the expression vector into an <u>E. coli</u>, yeast or animal cell. Examples of the monoclonal antibody-derived single chain antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 24 and VL contains the amino acid sequence of SEQ ID NO: 25. Examples of the human CDR-grafted antibody-derived disulfide stabilized antibody of the present invention include an antibody of which the VH contains the amino acid sequence of SEQ ID NO: 63.

A method for producing an anti-human IL-5R α chain monoclonal antibody which reacts specifically with a human IL-5R α chain or which inhibits the biological activity of human IL-5, and a method for producing an anti-human IL-5R α chain humanized antibody, an anti-human IL-5R α chain single chain antibody and an anti-human IL-5R α chain disulfide stabilized antibody all of which inhibit the biological activity of human IL-5, as well as a method for detecting and determining a human interleukin-5 receptor α chain by means of said antibodies will now be explained in detail.

1. Production of anti-hIL-5R α monoclonal antibody

(1) Preparation of antigen

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A cell having hIL-5R α expressed on the cell surface or a cell membrane fraction thereof, or an hIL-5R α -expressing cell CTLL-2 (h5 R) or a cell membrane fraction thereof can be used as an antigen for producing an anti-hIL-5R α monoclonal antibody. CTLL-2 (h5 R) is an hIL-5R α -expressing cell which was created by inserting a cDNA encoding a full length sequence of an pre-cloned hIL-5R α (J. Exp. Med., <u>175</u>, 341 (1992)) into an expression vector for animal cells such as pCAGGS (Gene, 108, 193 (1991)) and transfecting the expression vector into murine T cell line CTLL-2.

For expression in a prokaryotic host cell such as \underline{E} . \underline{coli} , a full length or partial fragment of cDNA encoding hIL-5R α can be inserted into an expression vector such as commercially available pGEX (Pharmacia), pET system (Novagen), pMKex1 to be described in section (11) of Example 1 below or the like and the full length hIL-5R α sequence or a partial

fragment thereof can be expressed either as such or as a fusion protein. After disruption of the cell, the protein expressed by \underline{E} . \underline{coli} can be purified by SDS-polyacrylamide electrophoresis, affinity chromatography based on the nature of the fusion protein, or the like.

In the method of expressing the full length IL-5R α sequence or a partial fragment thereof either as such or as a fusion protein, eukaryotic host cells such as insect cells, mammalian cells and the like can be used.

In the case of using a mammalian cell, a full length or a partial fragment of cDNA encoding hIL-5R α is inserted into a vector such as pAGE107 (Cytotechnology, $\underline{3}$, 133 (1990)), pAGE103 (J. Biochem., $\underline{101}$, 1307 (1987)), pAGE210 to be described in section (1) of Example 1 below or the like to thereby construct an expression vector for the protein. In order to express efficiently the full length hIL-5R α sequence encoded by the cDNA or a partial fragment thereof either as such or as a fused protein, the nucleotide sequence encoding a signal peptide in the cDNA is preferably replaced by the nucleotide sequence encoding a signal peptide of a protein which can be expressed at a high level in a eukaryotic host cell. Known signal peptides of proteins including those of human growth hormone, anti-ganglioside GD3 chimeric antibody KM871 (Kokai No. 304989/93) and the like are preferably used.

The thus constructed expression vector can be transfected into host cells by a known method such as electroporation (Kokai No. 257891/90; Cytotechnology, $\underline{3}$, 133 (1990)), lipofectin method (Proc. Natl. Acad. Sci., $\underline{84}$, 7413 (1987)) or the like. The cultivation of the cells in an appropriate medium can result in the production of the full length hIL-5R α sequence or a partial fragment thereof either as such or as a fusion protein in the cells or the culture supernatant. A serum-free medium is preferably used because it can facilitate the purification of the partial fragment or fusion protein of hIL-5R α produced in the culture supernatant.

In the case of using an insect cell, a full length or a partial fragment of cDNA encoding hIL-5R α is inserted using a Baculo Gold Starter Kit (Pharmingen) to prepare a recombinant baculovirus and insect c .lls of Sf9, Sf21 (Pharmingen) or the like are infected with the recombinant virus such that the full length hIL-5R α sequence or a partial fragment thereof is produced either as such or as a fusion protein in the cells or the culture supernatant (Bio/Technology, <u>6.</u> 47 (1988)).

The full length hIL-5R α sequence or a partial fragment or fusion protein thereof produced by the animal or insect cells can be purified from the culture supernatant or the like by a known method of protein purification such as salting-out, affinity chromatography ion-exchange chromatography or the like and can be used as an antigen. Particularly in the case where the hIL-5R α is produced as a fusion protein with a constant region of immunoglobulin, it is preferably purified using an affinity column having fixed thereto protein A which has specific affinity for the constant region of immunoglobulin.

(2) Immunization of animal and preparation of antibody-producing cells

Any animals such as mice, rats, hamsters, rabbits and the like can be used as animals to be immunized, provided that they can be used to prepare hybridomas. An embodiment in which mice or rats are used will be explained herein. Mice and rats of 3-20 weeks old are immunized with shlL-5R α or CTLL-2 which have hlL-5R α expressed on the surface (J. Exp. Med., 177, 1523 (1993)) as an antigen and antibody-producing cells are collected from the spleens, lymph nodes and peripheral bloods of the animals. Immunization is performed by administering the animals with the antigen together with an appropriate adjuvant such as complete Freund's adjuvant or a combination of aluminum hydroxide gel and pertussis vaccine either subcutaneously, intravenously or intraperitoneally. The antigen is administered 5-10 times at intervals of 1-2 weeks after the first administration. Blood is collected from the ophthal venous plexus at day 3-7 after each administration and the serum is examined for a reactivity with the antigen by enzyme immunoassay ("Enzyme Immunoassay (ELISA)", published by Igakushoin, 1976).

A mouse or rat whose serum shows a satisfactory antibody titer to shIL-5R α or the cells which have hIL-5R α expressed on the surface, which are used for immunization, can be used as a source of antibody-producing cells.

In order to perform fusion of a spleen cell with a myeloma cell, the spleen is removed from the immunized mouse at day 3-7 after the final administration of the antigenic substance and spleen cells are collected. The spleen is sliced in an MEM medium (Nissui Pharmaceuticals) and dispersed with a pair of tweezers. After centrifugation (1,200 rpm, 5 min), the supernatant is removed. The precipitate is treated with a Tris-ammonium chloride buffer (pH 7.65) for 1-2 minutes to remove erythrocytes and washed with MEM medium 3 times to prepare splenocytes for use in cell fusion.

(3) Preparation of myeloma cells

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An established cell line from a mouse or a rat is used as a myeloma cell. Examples include myeloma cell lines P3-X63Ag8-U1 (P3-U1) (Curr. Topics Microbiol. Immunol., $\underline{81}$, 1 (1978); Europ. J. Immunol., $\underline{6}$, 511 (1976)), SP2/0-Ag14 (SP-2) (Nature, $\underline{276}$, 269 (1978)), P3-X63-Ag8653 (653) (J. Immunol., $\underline{123}$, 1548 (1979)) and P3-X63-Ag8 (X63) (Nature, $\underline{256}$, 495 (1975)) which are derived from 8-azaguanine-tolerant mice (BALB/c). These cell lines can be subcultured in 8-azaguanine medium which is RPMI-1640 medium supplemented with glutamine (1.5 mM), 2-mercaptoethanol (5 x 10^{-5} M), gentamicin (10 μ g/ml) and fetal calf serum (FCS) (CSL, 10%) (hereinafter referred to as "normal

medium"), which is further supplemented with 8-azaguanine (15 μ g/ml). They should be subcultured in a normal medium 3-4 days before cell fusion to ensure a cell count of at least 2 x 10⁷ cells on the day of cell fusion.

(4) Cell fusion

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The antibody-producing cells described in 1 (2) and the myeloma cells described in 1 (3) are washed thoroughly with MEM medium or PBS (1.83 g of disodium phosphate, 0.21 g of monopotassium phosphate, 7.65 g of sodium chloride, 1 L of distilled water, pH 7.2). These cells are mixed such that a cell count ratio of the antibody-producing cells to the myeloma cells is 5-10:1. After centrifugation (1,200 rpm, 5 min), the supernatant is removed. The precipitated cells are dispersed and a mixed solution composed of ethylene glycol-1000 (PEG-1000)(2 g), MEM (2 ml) and dimethyl sulfoxide (DMSO) (0.7 ml) is then added to the cells in an amount of 0.2-1 ml/10⁸ antibody-producing cells while stirring. An MEM medium (1-2 ml) is added several times at intervals of 1-2 minutes and an additional MEM medium is then added such that the total volume is 50 ml. After centrifugation (900 rpm, 5 min), the supernatant is removed. The cells are dispersed gently and then suspended gently in 100 ml of a HAT medium (a normal medium supplemented with 10⁻⁴ M hypoxanthine, 1.5 x 10⁻⁵ M thymidine and 4 x 10⁻⁷ M aminopterin) by suction and blowoff with a pipette.

The cell suspension is dispensed in a 96-well culture plate in an amount of 100 μ l/well and cultured in a 5% CO₂ incubator at 37 °C for 7-14 days.

After the cultivation, an aliquot of the culture supernatant is examined by enzyme immunoassay to be described in 1 (5) to select a well that is reactive specifically with a recombinant protein such as a fusion protein with shIL-5R α or hIL-5R α described in 1 (1). Subsequently, cloning by limiting dilution is repeated twice. An aminopterin-free HAT medium is used in the first cloning and a normal medium in the second cloning. A cell exhibiting a high antibody titer stably is selected as a hybridoma cell line which produces a mouse or rat anti-hIL-5R α monoclonal antibody.

(5) Selection of mouse or rat anti-human IL-5R α monoclonal antibody

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A mouse or rat anti-hIL-5R α monoclonal antibody-producing hybridoma is selected in accordance with a method as described in Antibodies, A Laboratory Manual, Cold Spring Harbor Laboratory, Chapter 14 (1988) by the measurement method described below. By the method, the activity of an anti-hIL-5R α antibody in the culture supernatant of the transformants producing an anti-hIL-5R α humanized antibody, a single chain antibody or a disulfide stabilized antibody which are to be described below or the activities of all purified anti-hIL-5R α antibodies can be determined.

An appropriate plate is coated with shIL-5R α or a recombinant protein such as a fusion protein with hIL-5R α described in 1 (1). The plate is reacted with a primary antibody which is the hybridoma culture supernatant or the purified antibody to be obtained in 1 (6) and reacted with a secondary antibody which is an anti-mouse immunoglobulin antibody or an anti-rat immunoglobulin antibody which is labeled with biotin, an enzyme, an chemiluminescent substance or a radioactive compound. Subsequently, a reaction is performed in accordance with the specific kind of the label, whereby a hybridoma that is reactive specifically with hIL-5R α is selected as a hybridoma producing a mouse anti-hIL-5R α monoclonal antibody.

If the culture supernatant of the transformants producing an anti-hIL-5R α humanized antibody, a single chain antibody or a disulfide stabilized antibody, or an antibody purified therefrom is reacted as a primary antibody, an anti-human immunoglobulin antibody labeled with biotin, an enzyme, an chemiluminescent substance or a radioactive compound is used as a secondary antibody and a reaction is performed in - accordance with the specific kind of the label for detection.

An appropriate plate is coated with shIL-5R α or a recombinant protein such as a fusion protein with a recombinant protein hIL-5R α described in 1 (1). Any one of the hybridoma culture supernatant, the culture supernatant of the transformants producing an anti-hIL-5R α humanized antibody, a single chain antibody or a disulfide stabilized antibody, or an antibody purified therefrom is mixed and reacted with human IL-5 labeled with biotin, an enzyme, an chemiluminescent substance or a radioactive compound. Subsequently, a reaction is performed in accordance with the specific kind of the label so as to determine an activity in inhibiting the binding of human IL-5 to human IL-5R α . This method is used to screen hybridomas for selection of one having a high inhibitory activity against human IL-5.

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(6) Production of mouse or rat monoclonal antibody

A 8-10 week-old mouse or nude mouse is treated with pristane. More specifically, the mouse is administered intraperitoneally with pristane (2,6,10,14-tetramethylpentadecane, 0.5 ml) and bred for 2 weeks. The mouse is administered intraperitoneally with the mouse or rat anti-hIL-5R α monoclonal antibody-producing hybridoma cell lines (as obtained in 1 (3)) in an amount of 2 x 10⁷-5 x 10⁶ cells/mouse. The hybridoma caused ascites tumor after 10-21 days administration. The ascites is collected from the mouse and centrifuged (3,000 rpm, 5 min) to remove a solid portion. The precipitate is salted out and applied to a column for a caprylic acid precipitation, or a DEAE-Sepharose column, a protein A-column or a Cellulofine GSL2000 column (Biochemical Industry) to collect IgG or IgM fractions. These fractions are

used as a purified monoclonal antibody.

The subclass of the antibody is determined using a mouse or rat monoclonal antibody typing kit. The mass of the protein is calculated by a Lowry method or from the absorbance at 280 nm.

- 2. Production of anti-human IL-5R α humanized antibody
 - (1) Construction of humanized antibody expression vector

In order to produce a humanized antibody from a non-human animal antibody, a humanized antibody expression vector is prepared. The humanized antibody expression vector is a vector for expression in animal cells into which a gene encoding CH and CL, C regions of a human antibody, have been transfected. Such an expression vector is constructed by inserting two genes, one encoding CH of a human antibody and the other encoding CL of a human antibody, into an expression vector for animal cells. Any C regions of a human antibody such as C_{γ} 1 and C_{γ} 4 of a human antibody H chain, C_{κ} of a human antibody L chain and the like can be used. A chromosomal DNA consisting of an exon(s) and an intron(s) or cDNA can be used as a gene encoding a C region of a human antibody. Any expression vectors can be used as expression vectors for animal cells, provided that they can incorporate and express a gene encoding a C region of a human antibody. Examples are pAGE107 (Cytotechnology, 3, 133 (1990)), pAGE103 (J. Biochem., 101, 1307 (1987)), pHSG274 (Gene, 27, 223 (1984)), pKCR (Proc. Natl. Acad. Sci., 78, 1527 (1981)) and pSG1 β d2-4 (Cytotechnology, 4, 173 (1990)). A promoter and an enhancer to be used in preparation of an expression vector for animal cells are exemplified by an SV40 early promoter and enhancer (J. Bioche m., 101, 1307 (1987)), a Moloney mouse leukemia virus LTR promoter and enhancer (Biochem. Biophys. Res. Comun., 149, 960 (1987)), an immunoglobulin H chain promoter (Cell, 41, 479 (1985)) and enhancer (Cell, 33, 717 (1983)), and the like.

The humanized antibody expression vector may be either of a type in which a gene encoding an antibody H chain and a gene encoding an antibody L chain exist on separate vectors or of a type in which both genes exist on the same vector (tandem type). In terms of ease of construction of a humanized antibody expression vector, easiness of introduction into animal cells, balance between the expression amounts of antibody H and L chains in the animal cells and for other reasons, a tandem type of humanized antibody expression vector is more preferred (J. Immunol. Methods, 167, 271 (1994)).

(2) Preparation of cDNA encoding VH and VL of non-human animal antibody

cDNA encoding VH and VL of a non-human animal antibody such as a mouse anti-human IL-5R α chain monoclonal antibody is obtained, for example, as follows:

mRNA is extracted from an anti-human IL-5R α chain monoclonal antibody-producing cell such as a mouse anti-human IL-5R α chain antibody-producing hybridoma and used to synthesize cDNA. The synthesized cDNA is inserted into a vector such as a phage or a plasmid to prepare a cDNA library. From the library, with a portion in a V or C region of a non-human animal antibody such as a mouse antibody being used as a probe, a recombinant phage or plasmid which contains cDNA encoding VH and a recombinant phage or plasmid which contains cDNA encoding VL are isolated separately. The full nucleotide sequences of VH and VL of an antibody of interest which exist on the recombinant phage or plasmid are determined and the full amino acid sequences of the VH and VL are deduced from the nucleotide sequences.

(3) Construction of human chimeric antibody expression vector

A human chimeric antibody expression vector can be constructed by inserting cDNA encoding VH and VL of a non-human animal antibody in a region upstream of the gene encoding CH and CL of the human antibody on the humanized antibody expression vector which has been constructed in 2 (1). For example, a restriction enzyme recognition site for cloning of cDNA encoding VH and VL of a non-human animal antibody is created preliminarily in a region upstream of a gene encoding CH and CL of the human antibody on a chimeric antibody expression vector. At the cloning site, cDNA encoding a V region of a non-human animal antibody is inserted through a synthetic DNA (see below) to prepare a human chimeric antibody expression vector. The synthetic DNA consists of a nucleotide sequence at the 3' end of a V region of the non-human animal and a nucleotide sequence at the 5' end of a C region of the human antibody and are prepared by a DNA synthesizer such that it has appropriate restriction enzyme sites at both ends.

(4) Identification of CDR sequences of non-human animal antibody

VH and VL which form an antigen-binding site of an antibody consist of 3 complementarity determining regions (CDRs) having a wide variety of sequences which link the VH and VL to 4 framework regions (hereinafter referred to as "FR regions") having relatively conserved sequences (Sequences of Proteins of Immunological Interest, US Dept.

Health and Human Services, 1991). The amino acid sequence of the respective CDR (CDR sequence) can be identified by comparison with the amino acid sequences of V regions of known antibodies (Sequences of Proteins of Immunological Interest, US Dept. Health and Human Services, 1991).

(5) Construction of cDNA encoding V region of human CDR-grafted antibody

cDNA encoding VH and VL of a human CDR-grafted antibody can be obtained as follows:

In the first step, for each of VH and VL, the amino acid sequence of FR in a V region of a human antibody to which CDR in a V region of a non-human animal antibody of interest is to be grafted is selected. Any amino acid sequences of FRs in V regions derived from human antibodies can be used as the amino acid sequences of FRs in V regions of human antibodies. For example, the amino acid sequences of FRs in V regions of human antibodies recorded in Protein Data Bank and amino acid sequences common to subgroups of FRs in V regions of human antibodies (Sequences of Proteins of Immunological Interest, US Dept. Health and Human Services, 1991) can be used. In order to produce a human CDR-grafted antibody having an excellent activity, an amino acid sequence having high homology with the amino acid sequence of a V region of a non-human animal antibody of interest is desired. In the second step, a DNA sequence encoding the selected amino acid sequence of FR in a V region of a human antibody is ligated to a DNA sequence encoding the amino acid sequence of CDR in a V region of a non-human animal antibody and a DNA sequence encoding the amino acid sequences of VH and VL is designed. In order to obtain a DNA sequence designed to construct a CDR-grafted antibody variable region gene, several synthetic DNAs are designed for each strand such that the full DNA sequence is covered. Using the synthetic DNAs, polymerase chain reaction (hereinafter referred to as "PCR") is performed. For each strand, preferably 6 synthetic DNAs are designed in view of the reaction efficiency of PCR and the lengths of DNAs which can be synthesized. After the reaction, amplified fragments are subcloned into appropriate vectors and their nucleotide sequences are determined, thereby obtaining a plasmid which contains cDNA encoding the amino acid sequence of a V region of each strand of a human CDR-grafted antibody of interest. Alternatively, cDNA encoding the amino acid sequence of a V region of each strand of a human CDR-grafted antibody of interest may be constructed by synthesizing the full sequences of sense and antisense strand using synthetic DNAs consisting of about 100 bases and subjecting them to annealing and ligation.

(6) Modification of the amino acid sequence of V region of human CDR-grafted antibody

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It is known that if a human CDR-grafted antibody is prepared by simply grafting only CDR in a V region of a non-human animal antibody of interest between FRs in V in region of a human antibody, its activity is lower than that of the original non-human animal antibody (BIO/TECHNOLOGY, 9, 266 (1991)). Hence, among the amino acid sequences of FR in a V region of a human antibody, an amino acid residue which takes part in direct binding to an antigen, an amino acid residue which interacts with an amino acid residue in CDR, or an amino acid residue which may take part in the maintenance of the steric structure of an antibody is modified to an amino acid residue that is found in the original non-human animal antibody such that the activity of the human CDR-grafted antibody is increased. For efficient identification of the amino acid residue, the steric structure of an antibody is constructed and analyzed by X-ray crystallography, computer-modeling or the like. However, no method for producing a human CDR-grafted antibody which can be applied to any antibodies has yet been established and, therefore, various attempts must currently be made on a case-by-case basis.

The modification of the selected amino acid sequence of FR in a V region of a human antibody can be accomplished using various primers for mutation by PCR described in 2 (5). Amplified fragments obtained by the PCR are subcloned into appropriate vectors and their nucleotide sequences are determined, thereby obtaining a vector containing cDNA into which a mutation of interest has been introduced (hereinafter referred to as "amino acid sequence-replaced vector").

Alternatively, the modification of an amino acid sequence in a narrow region may be accomplished by a PCR-mutagenesis method using primers for mutation consisting of 20-35 bases. More specifically, a sense mutation primer and an antisense mutation primer which consist of 20-35 bases and which contain DNA sequences encoding the amino acid residue to be modified are synthesized and used to perform 2-step PCR using as a template a plasmid which contains cDNA encoding the amino acid sequence of a V region which is to be modified. The finally amplified fragments are subcloned into appropriate vectors and their nucleotide sequences are determined, thereby obtaining an amino acid sequence-modified vector containing cDNA into which a mutation of interest has been introduced.

(7) Construction of human CDR-grafted antibody expression vector

A human CDR-grafted antibody expression vector can be constructed by inserting the cDNA encoding VH and VL of the human CDR-grafted antibody obtained in 2 (5) and 2 (6) in a region upstream of the gene encoding CH and CL of the human antibody in the humanized antibody expression vector described in 2 (1). For example, if recognition sites

for appropriate enzymes are introduced at the ends of the 5' and 3' terminal synthetic DNAs during PCR for construction of cDNA encoding the amino acid sequences of VH and VL of the human CDR-grafted antibody, the cDNA can be inserted in a region upstream of a gene encoding a C region of a desired human antibody such that it is expressed in an appropriate form.

(8) Transient expression of humanized antibodies and evaluation of their activities

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In order to evaluate the activities of a wide variety of humanized antibodies efficiently, the human chimeric antibody expression vector described in 2 (3), and the human CDR-grafted antibody expression vector described in 2 (7) or their modified vectors may be transfected into COS-7 cells (ATCC CRL1651) and humanized antibodies expressed transiently (Methods in Nucleic Acids Res., CRC Press, p.283, 1991), followed by determination of their activities.

Examples of the method for transfecting the expression vector into a COS-7 cell include a DEAE-dextran method (Methods in Nucleic Acids Res., CRC Press, p.283, 1991), a lipofection method (Proc. Natl. Acad. Sci., <u>84</u>, 7413 (1987)) and the like.

After transfection of the vector, the activities of the humanized antibodies in the culture supernatant can be determined by the enzyme immunoassay (ELISA) described in 1 (5) and the like.

(9) Stable expression of humanized antibodies and evaluation of their activities

Transformants which produce a humanized antibody stably can be obtained by transfecting into appropriate host cells the human chimeric antibody expression vector described in 2 (3) and the human CDR-grafted antibody expression vector described in 2 (7).

Examples of the method for transfecting the expression vector into host cells include electroporation (Kokai No. 257891/90, Cytotechnology, 3, 133 (1990)) and the like.

Any cells can be used as host cells into which the humanized antibody expression vector is to be transfected, provided that they can express a humanized antibody. Examples are mouse SP2/0-Ag14 cell (ATCC CRL1581), mouse P3X63-Ag8.653 cell (ATCC CRL1580), CHO cells which are detective in dihydrofolate reductase gene (hereinafter referred to as "DHFR gene") (Proc. Natl. Acad. Sci.,77, 4216 (1980)) and rat YB2/3HL.P2.G11.16Ag.20 cell (ATCC CRL1662, hereinafter referred to as "YB2/0 cell").

After transfection of the vector, transformants which express a humanized antibody stably are selected in accordance with the method disclosed in Kokai No. 257891/90, using an RPMI1640 medium containing G418 and FCS. The humanized antibody can be produced and accumulated in a culture medium by culturing the selected transformants in a medium. The activity of the humanized antibody in the culture medium is determined by the method described in 1 (5) or the like. The production of the humanized antibody by the transformants can be increased by the method described in Kokai No. 257891/90, utilizing a DHFR gene-amplification system or the like.

The humanized antibody can be purified from the culture supernatant of the transformants by using a protein A column (Antibodies, A Laboratory Manual, Cold Spring Harbor, Chapter 8, 1988). Any other conventional methods for protein purification can be used. For example, the humanized antibody can be purified by a combination of gel filtration, ion-exchange chromatography, ultrafiltration and the like. The molecular weight of the H chain or L chain of the purified humanized antibody or the antibody molecule as a whole is determined by polyacrylamide gel electrophoresis (SDS-PAGE) (Nature, 227, 680, (1970)), western blotting (Antibodies, A Laboratory Manual, Cold Spring Harbor Laboratory, Chapter 12, 1988) and the like.

The reactivity of the purified humanized antibody and the inhibition activity of the humanized antibody against IL-5 can be determined by the method described in 1 (5).

(10) Method of use of humanized antibody

The humanized antibody of the present invention can bind specifically to a human IL-5R α chain, thereby inhibiting the biological activity of IL-5. Hence, the humanized antibody of the present invention is expected to inhibit the function of eosinophils which are controlled in differentiation and growth by IL-5. Accordingly, the humanized antibody of the present invention will be useful in the treatment of diseases where eosinophils are associated with their pathogenesis. Since almost all portions of the humanized antibody of the present invention are derived from the amino acid sequence of a human antibody, it is expected not only to exhibit immunogenicity in the human body but also to maintain its effect for a long period of time. The humanized antibody of the present invention can be used either alone or in combination with at least one pharmaceutically acceptable adjuvant. For example, the humanized antibody is dissolved in physiological saline or an aqueous solution of glucose, lactose, mannitol or the like to prepare a pharmaceutical composition. Alternatively, the humanized antibody is lyophilized by a conventional method and sodium chloride is added to prepare an injection in a powder form. If necessary, the present pharmaceutical composition may contain any additive that is well known in the field of pharmaceutical preparations such as a pharmaceutically acceptable salt and the like.

The present pharmaceutical composition can be administered to mammals including human at a dose of 0.1-20 mg/kg/day of the humanized antibody, which may vary depending on the age and conditions of the patient and the like. The administration is given once a day (single dose or continuous administration), 1-3 times a week or once every 2-3 weeks by intravenous injection.

3. Production of anti-human IL-5R α single chain antibody

(1) Construction of single chain antibody expression vector

A vector for expression of a single chain antibody of a non-human animal antibody or a single chain antibody of a human CDR-grafted antibody can be constructed by inserting into a single chain antibody expression vector the cDNAs encoding VH and VL of a non-human animal antibody or a human CDR-grafted antibody which are described in 2 (2), 2 (5) and 2 (6). Any expression vectors can be used as single chain antibody expression vectors, provided that they can incorporate and express the cDNAs encoding VH and VL of a non-human animal antibody or a human CDR-grafted antibody. Examples are pAGE107 (Cytotechnology, 3, 133 (1990)), pAGE103 (J. Biochem., 101, 1307 (1987)), pHSG274 (Gene, 27, 223 (1984)), pKCR (Proc. Natl. Acad. Sci., 78, 1527 (1981)) and pSG1 β d2-4 (Cytotechnology, 4. 173 (1990)). A host for use in expressing a single chain antibody can be selected from among E. coli, yeast and animal cells and the like. In this case, an expression vector which is compatible with the specific host should be selected. The single chain antibody can be secreted out of the cell and transported into the periplasm region or retained within the cell by inserting a cDNA encoding an appropriate signal peptide into the expression vector.

A single chain antibody expression vector into which the cDNA encoding a single chain antibody of interest has been inserted can be constructed by inserting the cDNA encoding a single chain antibody consisting of VH-L-VL or VL-L-VH (where L is a peptide linker) into the selected expression vector in a region downstream of an appropriate promoter and a signal peptide.

The cDNA encoding a single chain antibody can be obtained by linking a VH encoding cDNA to a VL encoding cDNA through a synthetic DNA encoding a peptide linker having recognition sites for appropriate restriction enzymes at both the ends. It is important to optimize the linker peptide such that its addition does not interfere with the binding of VH and VL to an antigen. For example, the linker described by Pantoliano et al. (Biochemistry, <u>30</u>, 10117 (1991)) and its modified versions may be used.

(2) Expression of single chain antibody and evaluation of its activity

A transformant which produces a single chain antibody of interest can be obtained by transfecting the single chain antibody expression vector constructed in 3 (1) into an approprate host cell by electroporation (Kokai No. 257891/90; Cytotechnology, 3, 133 (1990)) or the like. After transfection of the expression vector, the activity of the single chain antibody in the culture supernatant can be determined by the method described in 1 (5) or the like.

The collection and purification of the single chain antibody of the present invention can be accomplished by a combination of known techniques. For example, if the single chain antibody is secreted in a medium, it can be concentrated by ultrafiltration and its collection and purification can be then performed by antigen affinity chromatography or ion-exchange chromatography or gel filtration. If the single chain antibody is transported into the periplasm region of the host cell, it can be concentrated by ultrafiltration following the application of an osmotic shock and its collection and purification can be then performed by antigen affinity chromatography or ion-exchange chromatography or gel filtration. If the single chain antibody is insoluble and exists as a granule (i.e., inclusion body), its collection and purification can be performed by lysis of the cell, repeated centrifugation and washing for isolation of the granule, solubilization with guanidine-HCI, an operation for returning the structure of the single chain antibody to an active structure and the subsequent purification of an active molecule.

The activity of the purified single chain antibody can be determined by the method described in 1 (5) or the like.

(3) Method of using single chain antibody

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The single chain antibody of the present invention can bind specifically to a human IL-5R α chain, and inhibit the biological activity of IL-5. Hence, the single chain antibody of the present invention is expected to inhibit the function of eosinophils which are controlled in differentiation and growth by IL-5. Accordingly, the single chain antibody of the present invention will be useful in the treatment of diseases in which eosinophils are associated with the pathogenesis. The single chain antibody of the present invention can be used either alone or in combination with at least one pharmaceutically acceptable adjuvant. For example, the single chain antibody is dissolved in physiological saline or an aqueous solution of glucose, lactose, mannitol or the like to prepare a pharmaceutical composition. Alternatively, the single chain antibody is lyophilized by a conventional method and sodium chloride is added to prepare an injection in a powder form. If necessary, the present pharmaceutical composition may contain any additive that is well known in the field of phar-

maceutical preparations such as a pharmaceutically acceptable salt and the like.

The present pharmaceutical composition can be administered to mammals including human at a dose of 0.1-20 mg/kg/day of the signal chain antibody, which may vary depending on the age and conditions of the patient and the like. The administration is given once a day (single dose or continuous administration), 1-3 times a week or once every 2-3 weeks by intravenous injection.

- 4. Production of anti-human IL-5R α disulfide stabilized antibody
- (1) Production of disulfide stabilized antibody

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A disulfide stabilized antibody can be produced by a process comprising the steps of providing cDNAs encoding VH and VL of a non-human animal antibody or cDNAs encoding VH and VL of a human CDR-grafted antibody, modifying the DNA sequence which corresponds to a one-amino acid residue at an appropriate position in the respective cDNA with a DNA sequence corresponding to a cysteine residue, expressing the modified cDNAs and purifying the resultant peptide and then forming a disulfide bond. The modification of an amino acid residue to a cysteine residue can be performed by a mutagenesis method using PCR described in 2 (5).

A disulfide stabilized antibody H chain expression vector and a disulfide stabilized antibody L chain expression vector can be constructed by inserting the resulting cDNAs encoding the modified VH and modified VL into appropriate expression vectors. Any expression vectors can be used as disulfide stabilized antibody expression vectors, provided that they can incorporate and express cDNAs encoding a modified VH and a modified VL. For example, pAGE107 (Cytotechnology, $\underline{3}$, 133 (1990)), pAGE103 (J. Biochem., $\underline{101}$, 1307 (1987)), pHSG274 (Gene, $\underline{27}$, 223 (1984)), pKCR (Proc. Natl. Acad. Sci., $\underline{78}$, 1527 (1981)), pSG1 β d2-4 (Cytotechnology, $\underline{4}$. 173 (1990)) and the like can be used. A host used to express a disulfide stabilized antibody L chain expression vector and a disulfide stabilized antibody H chain expression vector for formation of a disulfide stabilized antibody can be selected from among \underline{E} . \underline{coli} , yeast and animal cells, and the like. In this case, an expression vector which is compatible with the specific host should be selected. The disulfide stabilized antibody can - be secreted out of the cell and transported into the periplasm region or retained within the cell by inserting a cDNA encoding an appropriate signal peptide into the expression vector.

(2) Expression of disulfide stabilized antibody and evaluation of its activity

A transformant which produces a disulfide stabilized antibody H chain or a disulfide stabilized antibody L chain of interest can be obtained by transfecting into a host cell the disulfide stabilized antibody H chain expression vector or the disulfide stabilized antibody L chain expression vector that were constructed in 4 (1) by electroporation (Kokai No. 257891/90; Cytotechnology, 3. 133 (1990)) or the like. After introduction of the expression vector, the expression of the disulfide stabilized antibody H chain or disulfide stabilized antibody L chain in the culture supernatant or the like can be confirmed by the method described in 1 (5).

The collection and purification of the disulfide stabilized antibody H chain or disulfide stabilized antibody L chain can be accomplished by combinations of known techniques. For example, if the disulfide stabilized antibody H chain or disulfide stabilized antibody L chain is secreted in a medium, they can be concentrated by ultrafiltration and their collection and purification can be then performed by various types of chromatography or gel filtration. If the disulfide stabilized antibody H chain or disulfide stabilized antibody L chain is transported into the periplasm region of the host cell, they can be concentrated by ultrafiltration after the application of an osmotic shock to the cell and their collection and purification can be then performed by various types of chromatography or gel filtration. If the disulfide stabilized antibody H chain or disulfide stabilized antibody L chain is insoluble and exists as a granule (i.e., inclusion body), their collection and purification can be performed by lysis of the cells, repeated centrifugation and washing for isolation of the granule, solubilization with guanidine-HCl and subsequent performance of various types of chromatography or gel filtration.

The purified disulfide stabilized antibody H chain and disulfide stabilized antibody L chain are mixed and subjected to a refolding procedure for deriving an active structure (Molecular Immunology, <u>32</u>, 249 (1995)), thereby forming a disulfide bond. Subsequently, the active disulfide stabilized antibody can be purified by antigen affinity chromatography or ion-exchange chromatography or gel filtration. The activity of the disulfide stabilized antibody can be determined by the method described in 1 (5) or the like.

(3) Method of use of disulfide stabilized antibody

The disulfide stabilized antibody of the present invention can bind specifically to a human IL-5R α chain, thereby inhibiting the biological activity of IL-5. Hence, the disulfide stabilized antibody of the present invention is expected to inhibit the function of eosinophils which are controlled in differentiation and growth by IL-5. Accordingly, the disulfide stabilized antibody of the present invention will be useful in the treatment of diseases in which eosinophils are associated with the pathogenesis. The disulfide stabilized antibody of the present invention can be used either alone or in combi-

nation with at least one pharmaceutically acceptable adjuvant. For example, the single chain antibody or disulfide stabilized antibody is dissolved in physiological saline or an aqueous solution of glucose, lactose, mannitol or the like to prepare a pharmaceutical composition. Alternatively, the disulfide stabilized antibody is lyophilized by a conventional method and sodium chloride is added to prepare an injection in a powder form. If necessary, the present pharmaceutical composition may contain any additive that is well known in the field of pharmaceutical preparations such as a pharmaceutically acceptable salt and the like.

The present pharmaceutical composition can be administered to mammals including human at a dose of 0.1-20 mg/kg/day of the disulfide stabilized antibody, which may vary depending on the age and conditions of the patient and the like. The administration is given once a day (single dose or continuous administration), 1-3 times a week or once every 2-3 weeks by intravenous injection.

- 5. Method for detection and determination of human interleukin-5 receptor α chain using anti-human IL-5R α antibody
- (1) Immunocyte staining using anti-human IL-5R α antibody

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When immunocytes are suspended cells, they are used as such in the following treatment. When immunocytes are adherent cells, they are detached with trypsin in EDTA and then used in the following treatment. The immunocytes are suspended in an immunocyte stain buffer (PBS containing 1% BAS, 0.02% EDTA and 0.05% sodium azide) or the like and dispensed in an amount of 1 x 10^5 -2 x 10^6 cells. The culture supernatant of the anti-human IL-5R α monoclonal antibody-producing hybridoma obtained in 1 (4), the culture supernatant of the anti-human IL-5R α humanized antibody transformant obtained in 2 (9) or the purified antibody obtained in 1 (6) or 2 (9), or the product obtained by labeling the purified antibody with an appropriate labeling substance (e.g., biotin) by a known method (KOUSOKOUTAIHOU (Methods for Enzymes and Antibodies), published by Gakusai Kikaku, 1985) and diluting the labeled antibody with an immunocyte stain buffer or a 10% animal serum-containing immunocyte stain buffer to a concentration of 0.1-50 µg/ml is dispensed in an amount of 20-500 µl and reacted on ice for 30 minutes. When the culture supernatant of the mouse anti-human IL-5R α monoclonal antibody-producing hybridoma obtained in 1 (4), the anti-human IL-5R α humanized antibody transformant obtained in 2 (9) or the purified antibody obtained in 1 (6) or 2 (9) has been reacted, the cells are washed with an immunocyte stain buffer after completion of the reaction and an immunocyte stain buffer containing about 0.1-50 µg/ml of an anti-mouse immunoglobulin antibody, anti-rat immunoglobulin antibody or anti-human immunoglobulin antibody which have been labeled with a fluorochrome such as FITC or phycoerythrin is dispensed in an amount of 50-500 µl, followed by reaction on ice for 30 minutes in the dark. When the biotin-labeled monoclonal antibody has been reacted, streptoavidin labeled with a fluorochrome such as FITC or phycoerythrin is dispensed in an amount of 50-500 µl and reaction is performed on ice for 30 minutes in the dark. When the monoclonal antibody labeled with a fluorochrome such as FITC or phycoerythrin has been reacted, an immunocyte stain buffer containing about 0.1-50 μg/ml of the monoclonal antibody is dispensed in an amount of 50-500 μl and reaction is performed on ice for 30 minutes in the dark. In each of these cases, the reaction mixture is washed thoroughly with an immunocyte stain buffer after the reaction and subjected to an analysis with a cell sorter.

(3) Test for inhibition of growth of human IL-5-dependent cells using anti-human IL-5R α antibody

In order to show the biological inhibition activity of the obtained anti-human IL-5R α antibody, the effect on the growth of human IL-5-dependent cells is examined using human IL-5 dependent cells. Examples of the evaluation method include incorporation of tritium-labeled thymidine into cells, color development methods using cell counting kits and the like. A color development method used in the present invention will now be explained.

CTLL-2 (h5R) cells (1 x 10^4) are suspended in a normal medium (50 μ l) and dispensed in a 96-well culture plate. To the plate are added 25 μ l of a solution of the purified antibody (0.01-50 μ g/ml) obtained in 1 (6) or 2 (9) and a normal medium containing 0.4-40 ng/ml of human IL-5 and the mixture is cultured in a 5% CO₂ incubator at 37°C for 24-72 hours. Subsequently, a cell counting kit solution is added at 10 μ l/well and the cultivation is continued in a 5% CO₂ incubator at 37°C for 4 hours. After completion of the cultivation, the absorbance at 450 nm is determined with a microwell plate reader Emax (Molecular Device) and the CTLL-2 (h5R) cell growth-inhibiting activity of the respective antibody is calculated.

(3) Suppression of survival of human eosinophils by anti-human IL-5R α antibody

Human polymorphonuclear leukocyte fractions which contain eosinophils are prepared from human peripheral blood with a commercially available corpuscle separation medium such as a polymorphprep (Nikomed) or a percoll (Pharmacia). The fractions are suspended in a normal medium and the resulting cells are dispensed in a 96, 48 or 24-well culture plate in an amount of $1 \times 10^6 - 1 \times 10^7$ cells/well, followed by addition of human IL-5 to a final concentration of 0.001-10 ng/ml. The culture supernatant of the anti-human IL-5R α monoclonal antibody-producing hybridoma

obtained in 1 (4) or the culture supernatant of the anti-human IL-5R α humanized antibody transformant obtained in 2 (9) or the purified antibody obtained in 1 (6) or 2 (9) is added and the mixture is cultured in a 5% CO $_2$ incubator at 37 °C for 2-5 days. After completion of the cultivation, a cell sample is prepared from each well and stained by May-Grünwald-Giemsa staining method (SENSHOKUHOU NO SUBETE (Techniques for Staining, published by Ishiyaku Shuppan Cor., Ltd., 1988) or the like and the percentage of eosinophils is determined. The absence or presence of the activity of the monoclonal antibody in suppressing the viability enhancement of IL-5-dependent human eosinophils is confirmed by comparing the percentage of eosinophils in the absence of the anti-human IL-5R α antibody with that in the presence of the anti-human IL-5R α antibody.

(4) Determination of shIL-5R α using monoclonal antibody

A plate is coated with 0.1-50 μ g/ml of the purified antibody obtained in 1 (6) or 2 (9) as a primary antibody. The coated plate is reacted with 0.1-10,000 ng/ml of the purified shlL-5R α obtained in 1 (1) or a sample such as human serum. The plate is washed thoroughly and then reacted with a secondary antibody which is an anti-human IL-5R α antibody recognizing an epitope other than that recognized by the anti-human IL-5R α antibody which was selected for use as the primary antibody from the purified antibodies obtained in 1 (6) or 2 (9). The secondary antibody was labeled with biotin, an enzyme, a chemiluminescent substance, a radioactive compound or the like prior to the reaction. Subsequently, a reaction is performed in accordance with the label. A calibration curve is constructed on the basis of the reactivity with the purified shlL-5R and the concentration of shlL-5R in the sample is calculated.

(5) Detection of shIL-5R α by Western blotting

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The purified shIL-5R α obtained in 1 (1) is subjected to SDS polyacrylamide electrophoresis (SDS-PAGE) and then blotted on a polyvinylidene difluoride membrane (hereinafter referred to as "PVDF membrane", Millipore). The PVDF membrane is immersed in PBS supplemented with 1-10% bovine serum albumin (BSA) and left to stand at 4°C overnight for blocking, followed by thorough washing with PBS containing 0.05% Tween. The PVDF membrane is immersed in the culture supernatant of the hybridoma obtained in 1 (5) or a solution of the purified antibody obtained in 1 (6) at room temperature for 2 hours and washed thoroughly with PBS containing 0.05% Tween. The PVDF membrane is immersed in a solution of an anti-mouse immunoglobulin antibody or anti-rat immunoglobulin antibody as a secondary antibody at room temperature for 1 hour and washed thoroughly with PBS containing 0.05% Tween. The secondary antibody was labeled preliminarily with biotin, an enzyme, a chemiluminescent substance, a radioactive compound or the like. After removing the washing solution completely, a reaction is performed in accordance with the label on the secondary antibody and a check is made for the reactivity with a protein which agrees in the molecular weight to the purified shIL-5R α .

(6) Immunoprecipitation of shIL-5R α

An anti-mouse immunoglobulin antibody or anti-rat immunoglobulin antibody is diluted 10-1000 folds with PBS or other buffer. The dilutions are dispensed in a 96-well ELISA plastic plate at 50-200 μ l/well and left to stand at 4 °C overnight or at room temperature for at least 2 hours, whereby they are adsorbed on the plate. The plate is washed with PBS. PBS containing 1-10% BSA and the like is dispensed in the plate at 300 μ l/well and left to stand at 4 °C overnight or at room temperature for at least 30 minutes to achieve blocking. The plate is washed with PBS. The culture supernatant of the hybridoma obtained in 1 (5) or a solution of the purified antibody obtained in 1 (6) (0.01-50 μ g/ml) is added at 50-200 μ l/well and left to stand at 4 °C overnight, thereby adsorbing the antibody on the plate. After the plate is washed, the shIL-5R α obtained in 1 (1) is diluted with PBS or the like containing 1% BSA to a concentration of 0.1-100 μ g/ml and the dilutions are dispensed at 50-200 μ l/well, followed by reaction at 4 °C overnight. After the plate is washed with PBS or the like containing 0.05% Tween, a 1× -5× sample buffer for SDS-PAGE is dispensed at 50-200 μ l/well and shaken at room temperature for at least 30 minutes. After optional dilution with PBS, the solution is added to each lane in an amount of 5-25 μ l and subjected to SDS-PAGE, followed by blotting on a PVDF membrane or the like by a conventional method. The PVDF membrane is subjected to western blotting as described in 5 (5), thereby detecting shIL-5R α .

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows steps for constructing plasmid pAGE210.

Figure 2 shows the restriction map of plasmid pCAGGS-h5R.25.

Figure 3 shows steps for constructing plasmid pAl234.

Figure 4 shows steps for constructing plasmid pAl230.

Figure 5 shows steps for constructing plasmid pAl282.

Figure 6 shows steps for constructing plasmids pAl283 and pAl285.

Figure 7 shows steps for constructing plasmids pAl284 and pAl289.

Figure 8 shows steps for constructing plasmids pAl294 and pAl295.

Figure 9 shows steps for constructing plasmids pAl299 and pAl301.

Figure 10 shows steps for constructing plasmid pAl292.

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Figure 11 shows steps for constructing plasmid pAl297.

Figure 12 shows steps for constructing plasmid pMKex1.

Figure 13 shows steps for constructing plasmid pAl263.

Figure 14 shows the binding reactivities of anti-human IL-5R α monoclonal antibody KM1257 and KM1259 with a human IL-5R α -human immunoglobulin constant region fusion protein in an enzyme immunoassay.

Figure 15 shows steps for constructing plasmid pBSA.

Figure 16 shows steps for constructing plasmid pBSAE.

Figure 17 shows steps for constructing plasmid pBSH-S.

Figure 18 shows steps for constructing plasmid pBSK-H.

15 Figure 19 shows steps for constructing plasmids pBSH-SA and pBSK-HA.

Figure 20 shows steps for constructing plasmids pBSH-SAE and pBSK-HAE.

Figure 21 shows steps for constructing plasmids pBSH-SAEE and pBSK-HAEE.

Figure 22 shows steps for constructing plasmid pBSK-HAEESa1.

Figure 23 shows steps for constructing plasmid pBSX-S.

20 Figure 24 shows steps for constructing plasmid pBSX-SA.

Figure 25 shows steps for constructing plasmid pBSSC.

Figure 26 shows steps for constructing plasmid pBSMo.

Figure 27 shows steps for constructing plasmid pBSMoS.

Figure 28 shows steps for constructing plasmid pChilgLA1S.

Figure 29 shows steps for constructing plasmid pMohCκ.

Figure 30 shows steps for constructing plasmid pBSMoSal.

Figure 31 shows steps for constructing plasmid pBSMoSalS.

Figure 32 shows steps for constructing plasmid pBShCy1.

Figure 33 shows steps for constructing plasmid pMohC γ 1.

Figure 34 shows steps for constructing plasmid pMoγ1SP.

Figure 35 shows steps for constructing plasmid pMoκγ1SP.

Figure 36 shows steps for constructing plasmid pKANTEX93.

Figure 37 shows steps for constructing plasmid pKANTEX1259H.

Figure 38 shows steps for constructing plasmid pKANTEX1259.

Figure 39 shows SDS-PAGE (on 4-15% gradient gel) electrophoresis patterns of anti-human IL-5R α chain human chimeric antibody KM1399. The left of the Figure shows the pattern of electrophoresis under non-reducing conditions and the right of the Figure under reducing conditions. On the left-hand side, M is a lane of high molecular weight markers and 1 is a lane of KM1399. On the right-hand side, M is a lane of low molecular weight markers and 1 is a lane of KM1399.

Figure 40 shows the inhibition activities of anti-human IL-5R α chain mouse antibody KM1259 and anti-human IL-5R α chain human chimeric antibody KM1399 against binding of human IL-5 to a human IL-5 α chain. The vertical axis of the graph plots the inhibition activity and the horizontal axis, the antibody concentration. \bullet refers to the activity of KM1259 and \bigcirc , the activity of KM1399.

Figure 41 shows steps for constructing plasmid pT1259.

Figure 42 shows the results of evaluation of activity on the basis of transient expression of an anti-human IL-5R α chain human chimeric antibody using plasmid pT1259. The vertical axis of the graph plots the inhibition activity against binding of human IL-5 to a human IL-5R α chain and the horizontal axis plots the dilution factor for the transient expression-culture supernatant.

Figure 43 shows steps for constructing plasmid phKM1259HV0.

Figure 44 shows steps for constructing plasmid phKM1259LV0.

Figure 45 shows steps for constructing plasmid pKANTEX1259HV0.

Figure 46 shows steps for constructing plasmid pKANTEX1259HV0LV0.

Figure 47 shows SDS-PAGE (on 4-15% gradient gel) electrophoresis patterns of anti-human IL-5R α chain human CDR-grafted antibody KM8397. The left of the Figure shows the pattern of electrophoresis under non-reducing conditions and the right of the Figure under reducing conditions. M is a lane of molecular weight markers and 1 is a lane of KM8397.

Figure 48 shows the activities of anti-human IL-5R α chain human chimeric antibody KM1399 and anti-human IL-5R α chain human CDR-grafted antibody KM8397 in binding to a human IL-5 α chain. The vertical axis of the graph plots the activity in binding to the human IL-5 α chain and the horizontal axis, an antibody concentration. \bullet refers to the

activity of KM1399 and O, the activity of KM8397.

Figure 49 shows the results of evaluation of the activities of various modified versions of anti-human IL-5R α chain human CDR-grafted - antibodies in transient expression-culture supernatants in inhibiting binding of human IL-5 to a human IL-5 α chain. The vertical axis of the graph plots the inhibitory activity and the horizontal axis indicates the names of samples. The inhibitory activity is expressed in relative terms, with the activity of chimeric antibody KM1399 taken as 100.

Figure 50 shows the activities of various modified versions of anti-human IL-5R α chain human CDR-grafted anti-bodies in binding to a human IL-5 α chain. The vertical axis of each graph plots the activity in binding to the human IL-5 α chain and the horizontal axis, the antibody concentration. In the upper graph, \bullet refers to the activity of KM1399; \bigcirc , HV.0LV.0; \blacksquare , HV.2LV.0; \square , HV.0LV.3; and \blacktriangle , HV.3LV.3. In the lower graph, \bullet refers to the activity of KM1399; \bigcirc , HV.0LV.0; \blacksquare , HV.3LV.0; \square , HV.0LV.4; \blacktriangle , HV.1LV.4, \triangle , HV.2LV.4; and X, H V.3LV.4.

Figure 51 shows steps for constructing plasmid pBShCγ4.

Figure 52 shows steps for constructing plasmids pKANTEX1259γ4 and pKANTEX1259HV3LV0γ4.

Figure 53 shows SDS-PAGE (on 4-15% gradient gel) electrophoresis patterns of anti-human IL-5R α chain human chimeric antibody KM7399 of a human antibody IgG4 subclass and human IL-5R α chain human CDR-grafted antibody KM9399 of a human antibody IgG4 subclass. The left of the Figure shows the pattern of electrophoresis under non-reducing conditions and the right of the Figure under reducing conditions. On the left-hand side, M is a lane of high molecular weight markers, 1 is a lane of KM9399 and 2 is a lane of KM7399. On the right-hand side, M is a lane of low molecular weight markers, 1 is a lane of KM9399 and 2 is a lane of KM7399.

Figure 54 shows the activity of anti-human IL-5R α chain human chimeric antibody KM1399 of a human antibody IgG1 subclass, human IL-5R α chain human chimeric antibody KM7399 of a human antibody IgG4 subclass, anti-human IL-5R α chain human CDR-grafted antibody KM8399 of a human antibody IgG1 subclass and anti-human IL-5R α chain human CDR-grafted antibody KM9399 of a human antibody IgG4 subclass in binding to a human IL-5 α chain. The vertical axis of the graph plots the activity of binding to the human IL-5 α chain and the horizontal axis, the antibody concentration. \bigcirc refers to the activity of KM1399; \blacksquare , KM7399; \square , KM8399; and \blacksquare , KM9399.

Figure 55 shows the results of flowcytometric analysis of the reactivities of anti-human IL-5R α monoclonal anti-bodies KM1257, KM1259, KM1399, KM1399, KM1399, KM1399 and KM1399 with a human IL-5R gene-transfected CTLL-2 cell.

Figure 56 shows the results of examination of the inhibitory action of anti-human IL-5R α monoclonal antibodies KM1257, KM1259, KM1486, KM1399, KM7399, KM8399 and KM9399 against IL-5-dependent growth of a human IL-5R gene-transfected CTLL-2 cell.

Figure 57 shows the results of flowcytometric analysis of the reactivity of anti-human IL-5R α monoclonal antibody KM1259 with human eosinophils.

Figure 58 shows the results of examination of inhibitory action of anti-human IL-5R α monoclonal antibodies KM1257, KM1259, KM1486, KM1399, KM7399, KM8399 and KM9399 for the survival of human eosinophils.

Figure 59 shows the results of evaluation of a soluble human IL-5R α quantitative determination system using anti-human IL-5R α monoclonal antibody KM1257 and biotin-labeled KM1259.

Figure 60 shows the results of detection of shIL-5R α by Western blotting using anti-human IL-5R α monoclonal antibodies KM1257, KM1259 and KM1486.

Figure 61 shows the results of immunoprecipitation of shIL-5R α using anti-human IL-5R α monoclonal antibodies KM1257, KM1259 and KM1486.

Best Mode for Carrying out the Invention

45 EXAMPLE 1

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- 1. Preparation of Antigens
- (1) Construction Expression Vector for Animal Cell pAGE210

Expression vector for animal cell, pAGE210, was constructed as described below using expression vectors for animal cell pAGE207 (Kokai No. 46841/94) and pAGE148 (Kokai No. 205694/94).

Three μ g of plasmid pAGE207 or pAGE148 was dissolved in 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM dithiothreitol (hereinafter referred to as "DTT"). To the resultant mixture, 10 units each of Clal and Kpnl (both manufactured by Takara Shuzo; unless otherwise indicated, the restriction enzymes used hereinbelow are those manufactured by Takara Shuzo) were added and reacted at 37 °C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.5 μ g of a 4.7 kb DNA fragment containing the SV40 early promoter and enhancer (hereinafter referred to as "P_{SE}"), a hygromycin resistance gene and an ampicillin resistance gene was recovered from pAGE207 and about 0.5 μ g of a 4.3 kb DNA fragment containing a

dihydrofolate reductase (hereinafter referred to as "dhfr") gene was recovered from pAGE148.

The Clal-KpnI fragment obtained from pAGE207 (50 ng) and the KpnI-Clal fragment obtained from pAGE148 (50 ng) were dissolved in 20 μ I of T4DNA ligase buffer [a buffer containing 66 mM Tris-HCl (pH 7.5), 6.6 mM magnesium chloride, 10 mM DTT and 0.1 mM adenosine triphosphate (hereinafter referred to as "ATP"]. To the resultant mixture, 200 units of T4DNA ligase (Takara Shuzo) was added and ligation was performed at 12°C for 16 hours. Using the prepared recombinant plasmid DNA, E. coli strain JM109 was transformed to thereby obtain plasmid pAGE210 shown in Fig. 1.

(2) Making shIL-5R α cDNA into a Cassette for the Construction of an shIL-5R α Expression Vector

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In order to construct an shIL-5R α expression vector, the modification of the 5' and 3' non-translational region of shIL-5R α cDNA and the introduction of a restriction enzyme recognition sequence were carried out using the PCR method [Maniatis et al. (eds.), Molecular Cloning, 14.2, Cold Spring Harbor Laboratory, 1989] according to the procedures described below.

Plasmid pCAGGS-h5R.25 is obtained by inserting shIL-5R α cDNA into the known plasmid pCAGGS [Gene, 108, 193 (1991)] as shown in Fig. 2 [J. Exp. Med., 175, 341 (1992)]. Three μg of this pCAGGS-h5R.25 was added to 30 μl of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT. Then, 10 units of EcoRl was added thereto, and reacted at 37 °C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μg of a 1.4 kb DNA fragment containing shIL-5R α cDNA was recovered.

Then, 1 ng of the DNA fragment obtained above was dissolved in 50 μl of PCR buffer [a buffer containing 50 mM potassium chloride, 10 mM Tris-HCl (pH 8.3), 1.5 mM magnesium chloride, 0.2 mM deoxyadenosine triphosphate (hereinafter referred to as "dATP"), 0.2 mM deoxyguanosine triphosphate (hereinafter referred to as "dGTP"), 0.2 mM deoxycytosine triphosphate (hereinafter referred to as "dCTP") and 0.2 mM deoxythymidine triphosphate (hereinafter referred to as " dTTP")]. To the resultant mixture, 50 pmol each of a synthetic DNA having the base sequence shown in SEQ ID NO: 1 and a synthetic DNA having the base sequence shown in SEQ ID NO: 2 [both synthesized with an automatic DNA synthesizer; Model 380A (Applied Biosystems Co., Ltd.)] and 1.6 units of Vent DNA polymerase (New England BioLabs, Inc.) were added and PCR was performed through 30 cycles under a series of conditions of 94 °C for 1 minute, 55 °C for 2 minutes and 72 °C for 3 minutes using a Perkin Elmer DNA thermal cycler (this was also used for the other PCR reactions). After the completion of the reaction, 2 µl of a buffer containing 100 mM Tris-HCl (pH 7.5), 100 mM magnesium chloride, 500 mM sodium chloride and 10 mM DTT, 8 μl of distilled water, and 10 units of HindIII were added to 10 µl of the reaction mixture and reacted at 37 °C for 4 hours. Then, DNA fragments were recovered from the reaction mixture by ethanol precipitation [Maniatis et al. (eds.), Molecular Cloning, E.10, Cold Spring Harbor Laboratory, 1989] and redissolved in 20 μl of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT. To the resultant mixture, 10 units of BamHI was added and reacted at 37 °C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μ g of a 1.0 kb DNA fragment

In a separate step, 3 μ g of plasmid pUC19 (Pharmacia Biotech) was dissolved in 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. Thereafter, DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT. To the resultant mixture, 10 units of BamHI was added and reacted at 37 °C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.5 μ g of the HindIII/BamHI fragment from pUC19 was recovered.

One hundred ng of the HindIII/BamHI fragment from pUC19 and 50 ng of shIL-5R α cDNA fragment were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12 °C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAI234 shown in Fig. 3.

(3) Construction of a Human Soluble IL-5R α Expression Vector

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An shIL-5R α expression vector, pAl230, was constructed as described below by ligating the HindIII-BamHI fragment from pAGE210 obtained in subsection (1) of Example 1 to the shIL-5R α cDNA-containing HindIII-BamHI fragment from pAl234 obtained in subsection (2) of Example 1.

Briefly, 3 μg of pAGE210 was added to 30 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μl of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.5 μg of a 9.0 kb DNA fragment was recovered.

Three μg of pAl234 was added to 30 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μl of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μg of a 1.0 kb DNA fragment was recovered.

Subsequently, 300 ng of the HindIII-BamHI fragment from pAGE210 and 50 ng of the HindIII-BamHI fragment from pAl234 were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl230 shown in Fig. 4.

(4) Modification of the Signal Sequence

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In order to produce shIL-5R α efficiently in animal cells, the signal sequence of the cDNA coding for shIL-5R α was modified according to the procedures described below by introducing an EcoRV recognition sequence into the cDNA at the 3' end of the signal sequence and subsequently replacing the original signal sequence with a signal sequence from a human growth hormone [Science, 205, 602 (1979)] or anti-ganglioside GD3 chimeric antibody KM871 (Kokai No. Hei 5-304989) using synthetic DNAs.

Briefly, 3 μg of plasmid pAl234 obtained in subsection (2) of Example 1 was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of Hindll was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHl was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μ g of a 1.0 kb DNA fragment was recovered.

In a separate step, 3 μ g of plasmid pUC19 was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of Hincll was added and reacted at 37°C for 4 hours. Then, DNA fragments were recovered from the reaction mixture by ethanol precipitation and about 0.5 μ g of a Hincll fragment from pUC19 was recovered.

About 1 ng of the DNA fragment obtained above was dissolved in 50 μ l of PCR buffer, to which 50 pmol each of a synthetic DNA having the base sequence shown in SEQ ID NO: 2 and a synthetic DNA having the base sequence shown in SEQ ID NO: 3 and 1.6 units of vent DNA polymerase were added. Then, PCR was performed through 30 cycles under a series of conditions of 94°C for 1 minute, 48 °C for 2 minutes and 72 °C for 3 minutes. Then, the reaction mixture was subjected to agarose gel electrophoresis, and 0.5 μ g of about 0.9 kb cDNA fragment coding for a portion of hIL-5R α was recovered. Fifty ng of this DNA and 100 ng of the HincII fragment from pUC19 were dissolved in 20 μ l of T4 ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl280 shown in Fig. 5. Three μ g of the thus obtained plasmid pAl280 was added to 30 μ l of a buffer containing 10 mM Tris-HCI (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of Xbal was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ l of a buffer containing 20 mM Tris-HCI (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.8 μ g of a 2.8 kb DNA fragment was recovered.

In a separate step, 3 μ g of plasmid pAl234 was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of Xbal was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHl was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.2 μ g of a 0.8 kb DNA fragment was recovered.

Subsequently, 200 ng of the Xbal-BamHI from pAl280 and 50 ng of the Xbal-BamHI from pAl234 were dissolved in 20 μ I of T4 ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl282 shown in Fig. 5. Three μ g of this plasmid pAl282 was added to 30 μ I of a buffer containing 50 mM Tris-HCI (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of EcoRV was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ I of a buffer containing 20 mM Tris-HCI (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μ g of a 0.9 kb DNA fragment was recovered.

One µg each of a synthetic DNA having the base sequence shown in SEQ ID NO: 4 and a synthetic DNA having

the base sequence shown in SEQ ID NO: 5 were dissolved in 10 μ l of distilled water. The resultant mixture was heated at 95 °C for 5 minutes and then cooled to room temperature over 30 minutes for annealing. A hundred ng of the Hindll-BamHI fragment from pUC19 obtained in subsection (2) of Example 1, 50 ng of the EcoRV-BamHI fragment from pAl282, and 50 ng of the synthetic DNAs having the base sequences shown in SEQ ID NOS. 4 and 5 which had been annealed as described above were dissolved in 20 μ l of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl283 shown in Fig. 6.

One μg each of a synthetic DNA having the base sequence shown in SEQ ID NO: 6 and a synthetic DNA having the base sequence shown in SEQ ID NO: 9 were dissolved in 10 μ l of distilled water. The resultant mixture was heated at 95 °C for 5 minutes and then cooled to room temperature over 30 minutes for annealing. To this reaction mixture, 2.5 μ l of a buffer containing 500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT and 1 mM EDTA, 2.5 μ l of 10 mM ATP solution, 9 μ l of distilled water and 5 units of T4 polynucleotide kinase (Takara Shuzo) were added, and phosporylation was performed at 37°C for 2 hours. Separately, 1 μ g each of a synthetic DNA having the base sequence shown in SEQ ID NO: 7 and a synthetic DNA having the base sequence shown in SEQ ID NO: 8 were dissolved in 10 μ l of distilled water. The resultant mixture was heated at 95 °C for 5 minutes and then cooled to room temperature over 30 minutes for annealing.

One hundred ng of the HindIII-BamHI fragment from pUC19, 50 ng of the EcoRV-BamHI fragment from pAl282, and 50 ng each of the synthetic DNAs as prepared above were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the thus prepared recombinant plasmid DNA, E. coli strain JM109 was transformed to thereby obtain plasmid pAl285 shown in Fig. 6.

(5) Construction of Signal Sequence-Modified shIL-5R α Expression Vectors

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Human soluble IL-5R α expression vectors, pAl284 and pAl289, were constructed as described below by ligating the HindIII-BamHI fragment from pAGE210 obtained in subsection (1) of Example 1 to the HindIII-BamHI fragment containing human soluble IL-5R α cDNA from pAl283 or pAl285 obtained in subsection (4) of Example 1.

Briefly, 3 μg each of pAl283 and pAl285 were added separately to 30 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μl of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 10 mM potassium chloride and 1 mM DTT, to which 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.3 μg of a 1.0 kb DNA fragment was recovered for each of the plasmids used.

Three hundred ng of the HindIII-BamHI fragment from pAGE210 and 50 ng of the HindIII-BamHI fragment from pAI283 or pAI285 were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmids pAI284 and pAI289 shown in Fig. 7.

(6) Preparation of a Fusion Protein Composed of Human IL-5R α and Human Immunoglobulin Constant Region

A fusion protein in which the extracellular region of human IL-5R α was linked to a human immunoglobulin constant region (hereinafter referred to as "Fc") through a linker having an amino acid sequence of (Gly-Ser-Gly)₄ (hereinafter, this fusion protein is referred to as "hIL-5R α -Fc") was prepared according to the procedures described below.

As a cDNA coding for a human immunoglobulin constant region, the portion of the human chimeric antibody H chain expression vector pChilgHB2 (Kokai No. Hei 5-304989) which coded for the human lgG1 constant region was used. First, about 1 ng of pChilgHB2 was dissolved in 50 μ l of PCR buffer. To this solution, 50 pmol each of a synthetic DNA having the base sequence shown in SEQ ID NO: 10 and a synthetic DNA having the base sequence shown in SEQ ID NO: 11 and 1.6 units of vent DNA polymerase were added. Then, PCR was performed through 30 cycles under a series of conditions of 94°C for 1 minute, 48 °C for 2 minutes and 72 °C for 3 minutes. After the completion of the reaction, 2.5 μ l of a buffer containing 200 mM Tris-HCl (pH 8.5), 100 mM magnesium chloride, 1000 mM potassium chloride and 10 mM, 2.5 μ l of distilled water, and 10 units of BamHI were added to 20 μ l of the reaction mixture and reacted at 37 °C for 4 hours. After the completion of the reaction, the reaction mixture was subjected to agarose gel electrophoresis, and about 0.5 μ g of a 0.7 kb DNA fragment containing a cDNA coding for the human IgG1 constant region was recovered.

About 1 ng of pAl283 obtained in subsection (4) of Example 1 was dissolved in 50 µl of PCR buffer, to which 50 pmol each of a synthetic DNA having the base sequence shown in SEQ ID NO: 12 and a synthetic DNA having the base sequence shown in SEQ ID NO: 13 and 1.6 units of vent DNA polymerase were added. Then, PCR was performed through 30 cycles under a series of conditions of 94°C for 1 minute, 48 °C for 2 minutes and 72 °C for 3 minutes. After the completion of the reaction, 2.5 µl of a buffer containing 100 mM Tris-HCl (pH 7.5), 100 mM magnesium chloride,

500 mM sodium chloride and 10 mM DTT, 2.5 μ l of distilled water, and 10 units of HindIII were added to 20 μ l of the reaction mixture and reacted at 37 °C for 4 hours. After the completion of the reaction, the reaction mixture was subjected to agarose gel electrophoresis. Thereafter, about 0.5 μ g of a 1.0 kb DNA fragment containing a cDNA coding for the extracellular region of hIL-5R α was recovered.

Fifty ng or the 0.7 kb DNA fragment containing the cDNA coding for the human lgG1 constant region, 50 ng of the DNA fragment containing the cDNA coding for the extracellular region of hIL-5R α and 100 ng of the HindIII-BamHI fragment from pUC19 were dissolved in 20 μ l of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl294 shown in Fig. 8.

In a separate step, PCR reaction was conducted under conditions similar to those described above using pAl285 obtained in subsection (4) of Example 1 as a template and also using synthetic DNAs having the base sequences shown in SEQ ID NOS: 13 and 14, as primers. After the completion of the reaction, the reaction mixture was subjected to agarose gel electrophoresis. Subsequently, about 0.5 μ g of a 1.0 kb DNA fragment containing the cDNA coding for the extracellular region of human IL-5R α was recovered. Fifty ng of the thus obtained DNA fragment, 50 ng of the 0.7 kb DNA fragment containing the cDNA coding for the human IgG1 constant region and 100 ng of the HindIII-BamHI fragment from pUC19 were dissolved in 20 μ l of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAl295 shown in Fig. 8.

(7) Construction of a Fusion Protein Expression Vector

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An hIL-5R α -Fc expression vector, pAl299, was constructed as described below by ligating the HindIII-BamHI fragment from pAGE210 obtained in subsection (1) of Example 1 to the HindIII-BamHI fragment from pAl294 obtained in subsection (6) of Example 1 containing the cDNA coding for hIL-5R α -Fc.

Briefly, 3 μ g of plasmid pAl294 was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and redissolved in 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 100 mM potassium chloride and 1 mM DTT. To the resultant mixture, 10 units of BamHI was added and reacted at 37°C for 4 hours. After the reaction mixture was subjected to agarose gel electrophoresis, about 0.4 μ g of a 1.7 kb DNA fragment containing a cDNA coding for a fusion protein composed of human IL-5R α and the human immunoglobulin constant region was recovered.

One hundred ng of the HindIII-BamHI fragment from pAGE210 and 50 ng of the HindIII-BamHI fragment from pAI294 were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmid pAI299 shown in Fig. 9.

Further, an hIL-5R α -Fc expression vector, pAl301, was constructed similarly by ligating the HindIII-BamHI fragment from pAGE210 to the HindIII-BamHI fragment from pAl295 obtained in subsection (6) of Example 1 containing the cDNA coding for hIL-5R α -Fc.

$^{\circ}$ (8) Preparation of a Recombinant Virus for Expressing shIL-5R lpha in Insect Cells

For the production of a protein in insect cells, a recombinant virus inserting a gene of interest is prepared. The preparation of such a virus is performed through a process in which a cDNA coding for a gene of interest is incorporated into a special plasmid called "a transfer vector" and a subsequent process in which a wild-type virus and the transfer vector are co-transfected into insect cells to obtain a recombinant virus by homologous recombination. The processes described above were performed using BaculoGold Starter Kit (Cat. No. PM-21001K) manufactured by Pharmingen according to the manufacturer's manual.

Briefly, 3 μ g of pAl285 obtained in subsection (4) of Example 1 or pAl294 obtained in subsection (6) of Example 1 was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of HindIII was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and dissolved in 20 μ l of DNA polymerase I buffer [a buffer containing 5 mM Tris-HCl (pH 7.5), 1 mM magnesium sulfate, 0.01 mM DTT, 5 μ g/ml bovine serum albumin, 0.08 mM dATP, 0.08 mM dGTP, 0.08 mM dCTP and 0.08 mM dTTP]. To the resultant mixture, 5 units of E. coli DNA polymerase I Klenow fragment (Takara Shuzo) was added and reacted at 22 °C for 30 minutes, whereby the 5' sticky ends generated by the HindIII digestion were changed to blunt ends. Further, the reaction mixture was subjected to phenol-chloroform extraction followed by ethanol precipitation. To the precipitate, 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 100 mM potassium chloride and 1 mM DTT, and 10 units of BamHI were added and reacted at 37°C for 4 hours. The reaction mixture was subjected to agarose gel electrophoresis, and about 0.3 μ g of an approx. 1.0 kb DNA fragment containing the cDNA coding for shIL-5R α and about 0.3 μ g of a 1.7 kb DNA fragment containing

the cDNA coding for the fusion protein composed of human IL-5 α and the human immunoglobulin constant region were recovered.

Subsequently, 3 μ g of plasmid pVL1393 contained in BaculoGold Starter Kit (Pharmingen) was added to 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of EcoRl was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and dissolved in 20 μ l of DNA polymerase I buffer, to which 5 units of E. coli DNA polymerase I Klenow fragment was added and reacted at 22°C for 30 minutes, whereby the 5' sticky ends generated by the EcoRl digestion were changed to blunt ends. Further, the reaction mixture was subjected to phenol-chloroform extraction followed by ethanol precipitation. To the precipitate, 30 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, and 10 units of BgIII were added and reacted at 37°C for 4 hours. The reaction mixture was subjected to agarose gel electrophoresis, and about 0.9 μ g of an approx. 9.6 kb DNA fragment was recovered.

Thereafter, 200 ng of the thus obtained EcoRI (blunt end)-BgIII fragment from pVL1393 and 50 ng of the HindIII (blunt end)-BamHI fragment from pAl285 or pAl294 were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the recombinant plasmid DNA thus prepared, E. coli strain JM109 was transformed to thereby obtain plasmids pAl292 and pAl297 shown in Figs. 10 and 11, respectively.

The subsequent preparation of a recombinant virus was performed as described below by transfecting into an insect cell, Sf9 (obtained from Pharmingen), cultured in TMN-FH Insect Medium (Pharmingen), a linear baculovirus DNA (BaculoGold baculovirus DNA; Pharmingen) and the prepared transfer vector DNA by the lipofectin method [TAN-PAKUSHITSU, KAKUSAN, KOHSO (Protein, Nucleic Acid, Enzyme), <u>37</u>, 2701 (1992)].

Briefly, 1 μ g of pAl292 or pAl297 and 20 ng of the linear baculovirus DNA were dissolved in 12 μ l of distilled water, to which a mixture of 6 μ l of lipofectin and 6 μ l of distilled water was added and left at room temperature for 15 minutes. In a separate step, 1×10^6 Sf9 cells were suspended in 2 ml of Sf900-II medium (Gibco) and put in a plastic cell culture dish 35 mm in diameter. To this dish, a total volume of the above-described mixture of plasmid DNA, linear baculovirus DNA and lipofectin was added, and cells were cultured at 27 °C for 3 days. Thereafter, 1 ml of the culture supernatant containing a recombinant virus was taken. One ml of a fresh Sf900-II medium was added to the dish and cells were cultured at 27 °C for another 3 days. Then, an additional 1.5 ml of the culture supernatant containing a recombinant virus was obtained.

Subsequently, the thus obtained recombinant virus was propagated for the purpose of use in protein expression, according to the procedures described below.

Briefly, 2x10⁷ Sf9 cells were suspended in 10 ml of Sf900-II medium, put in a 175 cm² flask (Greiner) and left at room temperature for 1 hour to allow cells to adhere to the flask. Thereafter, the supernatant was removed, and 15 ml of a fresh TMN-FH Insect Medium and 1 ml of the above-obtained culture supernatant containing the recombinant virus were added to the flask. Then, cells were cultured at 27°C for 3 days. After the cultivation, the supernatant was centrifuged at 1,500xg for 10 minutes to remove cells. Thus, a viral solution to be used for protein expression was obtained.

With respect to the thus obtained solution of the recombinant virus, the viral titer was calculated by the method described below (BaculoGold Starter Kit Manual; Pharmingen). A number (6x10⁶) of Sf9 cells were suspended in 4 ml of Sf900-II medium, put in a plastic cell culture dish 60 mm in diameter and left at room temperature for 1 hour to allow cells to adhere to the dish. After the removal of the supernatant, 400 µl of a fresh Sf900-II medium and the above-described recombinant virus solution diluted 10,000 folds with Sf900-II medium were added to the dish and left at room temperature for 1 hour. Then, the medium was removed, and 5 ml of a medium containing 1% low melting point agarose (Agarplaque Agarose; Pharmingen) (a medium obtainable by mixing 1 ml of sterilized 5% aqueous Agarplaqueplus Agarose solution and 4 ml of TMN-FH Insect Medium and keeping the mixture at 42°C] was poured into the dish. After the dish was left at room temperature for 15 minutes, vinyl tape was wound round the dish to prevent dryness. Then, the dish was placed in an airtight plastic container and cells were cultured at 27°C for 6 days. After 1 ml of PBS containing 0.01% Neutral Red was added to the dish and cells were cultured for an additional day, the number of plaques formed was counted. From the operations described above, it was found that each of the recombinant virus solutions contained about 1x10⁷ plaque forming units(PFU)/ml of virus.

(9) Expression of shIL-5R α or hIL-5R α -Fc in Animal Cells

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The introduction of a plasmid into animal cells was performed according to the method of Miyaji et al. using electroporation [Cytotechnology, 3, 133 (1990)].

Briefly, 4 μ g of pAl289 obtained in subsection (5) of Example 1 or pAl301 obtained in subsection (7) of Example 1 was transfected into $4x10^6$ dhfr gene-deficient CHO cells [Proc. Natl. Acad. Sci., $\underline{77}$, 4216 (1980)], which were then suspended in 40 ml of RPMI1640-FCS(10) [RPMI1640 medium containing 10% FCS, 1/40 volume 7.5% NaHCO₃, 3% 200 mM L-glutamine solution (Gibco) and 0.5% penicillin/streptomycin solution (Gibco; containing 5000 units/ml penicillin and 5000 μ g/ml streptomycin); manufactured by Nissui Pharmaceuticals] and dispensed into a 96-well microtiter plate

(200 μ l/well). After the cells were cultured in a CO₂ incubator at 37°C for 24 hours, hygromycin (Gibco) was added to give a concentration of 0.5 mg/ml. Then, the cells were cultured for an additional 1-2 weeks. Cells were recovered from those wells which became confluent with the appearance of colonies of transformant, and suspended in RPMI1640-FCS(10) medium containing 0.5 mg/ml hygromycin and 50 nM methotrexate (hereinafter referred to as " MTX") to give a cell density of 1-2 x 10^5 cells/ml. The cell suspension was dispensed into a 24-well plate (2 ml/well) and the cells were cultured in a CO₂ incubator at 37°C for 1-2 weeks to thereby induce 50 nM MTX resistant clones.

The thus obtained 50 nM MTX resistant clones were suspended in RPMI1640-FCS(10) medium containing 0.5 mg/ml hygromycin and 200 nM MTX to give a cell density of 1-2 x 10^5 cells/ml. The cell suspension was dispensed into a 24-well plate (2 ml/well) and the cells were cultured in a $\rm CO_2$ incubator at 37 °C for 1-2 weeks to thereby induce 200 nM MTX resistant clones.

Further, the thus obtained 200 nM MTX resistant clones were suspended in RPMI1640-FCS(10) medium containing 0.5 mg/ml hygromycin and 500 nM MTX to give a cell density of 1-2 x 10^5 cells/ml. The cell suspension was dispensed into a 24-well plate (2 ml/well) and the cells were cultured in a CO₂ incubator at 37°C for 1-2 weeks to thereby induce 500 nM MTX resistant clones.

The above transformants were suspended in a serum-free medium for CHO cells, CHO-S-SFMII medium (Gibco), to give a cell density of 1-2 x 10^5 cells/ml, and the cell suspension was dispensed into 225 cm² flasks (Greiner) in an amount of 100 ml/flask. The cells were cultured in a CO₂ incubator at 37 °C for 5-7 days and the culture medium was recovered when confluence was attained.

The purification of hIL-5R α from the culture supernatant was performed as follows. To 1 liter of the culture medium of pAl289-derived transformant, 29.2 g of sodium chloride and 20 ml of 1 M Tris-HCl (pH 7.4) were added. Then, the pH of the resultant mixture was adjusted to 7.4 with 1 N sodium hydroxide solution. A column was packed with about 10 ml of Concanavalin A-Sepharose (Pharmacia) gel and then washed with 50 ml of a buffer containing 20 mM Tris-HCl (pH 7.4) and 0.5 M sodium chloride at a flow rate of 0.5 ml/min. After the washing, the mixture containing shIL-5R α prepared as described above was applied to the Concanavalin A-Sepharose column at a flow rate of 0.5 ml/min. Then, the column was washed with 80 ml of a buffer containing 20 mM Tris-HCl (pH 7.4) and 0.5 M sodium chloride at a flow rate of 0.5 ml/min. Thereafter, the protein adsorbed on Concanavalin A-Sepharose was eluted and, simultaneously, the eluate was fractionated into 1 ml fractions (fractions 1-30) with 15 ml of a buffer containing 20 mM Tris-HCI (pH 7.4) and 0.5 M sodium chloride and 15 ml of a buffer containing 0.5 M α -methylmannoside, 20 mM Tris-HCl (pH 7.4) and 0.5 M sodium chloride by linearly changing the α-methylmannoside concentration from 0 to 0.5 M. Further, 20 ml of a buffer containing 1 M α -methylmannoside, 20 mM Tris-HCl (pH 7.4) and 0.5 M sodium chloride was applied to the column and the eluate was fractionated into 2 ml fractions (fractions 31-40). The protein concentration of each fraction was measured using a protein concentration measurement kit (Bio-rad) and fractions 10-40 having high protein concentration were recovered. The resultant protein solution was concentrated by a factor of about 10 using Centricon-30 (Amicon), placed in a dialysis tube and dialyzed against PBS. Thus, a purified shIL-5R α (protein concentration: 4 mg/ml; 3.5 ml) was obtained.

In a separate step, hIL-5R α -Fc was obtained as follows. A column was packed with about 5 ml of Protein A-Sepharose gel and then washed with 50 ml of PBS. After the washing, 1 liter of the culture medium of the pAl301-derived transformants described above was applied to the Protein A-Sepharose column at a flow rate of 0.5 ml/min. Then, the column was washed with 50 ml of PBS. Thereafter, 20 ml of 0.1 M citrate buffer (pH 3.0) was applied to the column to thereby elute the protein adsorbed on Protein A-Sepharose and, simultaneously, fractionate the eluate into 1-ml fractions. To each of the fractions, 0.15 ml of 2M Tris-HCl (pH 9.0) was added for pH adjustment. The protein concentration of each fraction was measured using a protein concentration measurement kit (Bio-rad) and those fractions having high protein concentration were recovered. The resultant protein solution was placed in a dialysis tube and dialyzed against PBS. Thus, a purified hIL-5R α -Fc (protein concentration: 1.8 mg/ml; 5.5 ml) was obtained.

(10) Expression of shIL-5R α or hIL-5R α -Fc in Insect Cells

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The expression of shIL-5R α and hIL-5R α -Fc was performed by the procedures described below according to the manual attached to BaculoGold Starter Kit (Pharmingen).

The recovery of shIL-5R α and hIL-5R α -Fc from culture mediums was performed using Concanavalin A-Sepharose and Diethylaminoethyl(DE AE)-Sepharose, or Protein A-Sepharose (all manufactured by Pharmacia Biotech), respectively.

shIL-5R α was obtained as follows. Briefly, $6x10^6$ Sf9 cells were suspended in 45 ml of Grace's Insect Medium (Gibco) containing 10% FCS in a 225 cm² flask (Greiner) and cultured at 27 °C for 3-4 days. After the culture supernatant was removed, 30 ml of a fresh Grace's Insect Medium containing 10% FCS and 1 ml of a solution in which the recombinant virus derived from the transfer vector pAl292 obtained in 1(8) of Example 1 was contained at a concentration of approx. $1x10^7$ PFU/ml were added. The cells were cultured at 27 °C for one additional day. Then, after the removal of the culture supernatant, 45 ml of a fresh Sf900-II medium was added and the cells were cultured for 2-3 days. After the completion of the cultivation, the culture supernatant was recovered and centrifuged at 1,500xg for 10

minutes, to thereby obtain a supernatant. To the resultant culture medium, sodium chloride was added to give a final concentration of 0.5 M. Then, 1/50 volume of 1 M Tris-HCl (pH 7.4) was added and the pH of the resultant mixture was adjusted to 7.4 with 1 N sodium hydroxide solution.

A column was packed with about 10 ml of Concanavalin A-Sepharose gel and washed with 50 ml of a buffer containing 20 mM Tris-HCl (pH 7.4) and 0.5 mM sodium chloride at a flow rate of 0.5 ml/min. After the washing, 500 ml of the shlL-5R α containing culture medium prepared as described above was applied to the Concanavalin A-Sepharose column at a flow rate of 0.5 ml/min. Then, the column was washed with 80 ml of a buffer containing 20 mM Tris-HCl (pH 7.4) and 0.5 mM sodium chloride at a flow rate of 0.5 ml/min. Thereafter, 60 ml of a buffer containing 1 M α -methylmannoside, 20 mM Tris-HCl (pH 7.4) and 0.5 M sodium chloride was applied to the column to thereby elute the protein adsorbed on Concanavalin A-Sepharose and, simultaneously, fractionate the eluate into 2-ml fractions. The protein concentration of each fraction was measured using a protein concentration measurement kit (Bio-rad). Those fractions with high protein-concentration were recovered in a total amount of 44 ml and dialyzed against 20 mM Tris-HCl (pH 7.4). Further, similar operations were performed on 900 ml of the shlL-5R α containing culture medium prepared as described above so as to recover those fractions with high protein-concentration in a total amount of 40 ml, which were dialyzed against 20 mM Tris-HCl (pH 7.4).

After the dialysis, the two protein solutions were combined and applied to a column packed with 10 ml of Diethylaminoethyl(DEAE)-Sepharose gel to have the protein adsorbed. The elution of shIL-5R α from the column was performed by linearly changing the sodium chloride concentration from 0 to 0.5 M. Thus, those fractions with high concentration of shIL-5R α were recovered in a total amount of 4 ml. This protein solution was placed in a dialysis tube and dialyzed against PBS. Thus, a purified shIL-5R α (protein concentration: 400 μ g/ml; 4.5 ml) was obtained.

In a separete step, hIL-5R α -Fc was obtained as follows. Briefly, $6x10^6$ Sf9 cells were suspended in 45 ml of Grace's Insect Medium (Gibco) containing 10% FCS in a 225 cm² flask (Greiner) and cultured at 27°C for 3-4 days. After the culture supernatant was removed, 30 ml of a fresh Grace's Insect Medium containing 10% FCS and 1 ml of a solution in which the recombinant virus derived from the transfer vector pAl297 obtained in 1(8) of Example 1 was contained at a concentration of approx. $1x10^7$ PFU/ml were added. The cells were cultured further at 27°C for one additional day. Then, after the removal of the culture supernatant, 45 ml of a fresh Sf900-II medium was added and the cells were cultured for 2-3 days. After the completion of the cultivation, the culture supernatant was recovered and centrifuged at 1,500xg for 10 minutes, to thereby obtain a supernatant.

A column was packed with about 5 ml of Protein A-Sepharose gel and washed with 50 ml of PBS. After the washing, 450 ml of the shIL-5R α -Fc containing culture medium as described above was applied to the Protein A-Sepharose column at a flow rate of 0.5 ml/min. Then, the column was washed with 50 ml of PBS. Thereafter, 20 ml of 0.1 M citrate buffer (pH 3.0) was applied to the column to thereby elute the protein adsorbed on Protein A-Sepharose and, simultaneously, fractionate the eluate into 1-ml fractions. To each of the fractions, 0.15 ml of 2 M Tris-HCl (pH 9.0) was added for pH adjustment. The protein concentration of each fraction was measured using a protein concentration measurement kit (Bio-rad) and those fractions with high protein concentration were recovered. The thus obtained protein solution was concentrated by a factor of about 3 using Centricon-30 (Amicon), placed in a dialysis tube and dialyzed against PBS. Thus, a purified shIL-5R α -Fc (protein concentration: 0.4 mg/ml; 1.8 ml) was obtained.

(11) Expression of an shIL-5R α Partial Fragment in E. coli

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The expression of an shIL-5R α partial fragment in E. coli was performed by inserting a DNA fragment containing a cDNA coding for an shIL-5R α partial fragment into E. coli expression vector pMKex1 to be described below so as to construct pAl263 and transform E. coli with pAl263.

Briefly, 3 μ g of plasmid pGHA2 (Kokai No. Sho 60-221091) was added to 30 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of EcoRl was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation. To these DNA fragments, 30 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, and 10 units of Clal were added and reacted at 37 °C for 4 hours. The reaction mixture was subjected to agarose gel electrophoresis, and about 0.3 μ g of the EcoRl-Clal fragment from pGHA2 containing the promoter region was recovered.

Three μ g of plasmid pTerm2 (Kokai No. 227075/90) was added to 30 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of EcoRl was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation. To these DNA fragments, 30 μ l of a buffer containing 10 mM Tris-HCl (pH 8.4), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT and 10 units of Nsil were added and reacted at 37°C for 4 hours. The reaction mixture was subjected to agarose gel electrophoresis, and about 0.8 μ g of the EcoRl-Nsil fragment from pTerm2 was recovered.

Fifty ng of the EcoRI/Clal fragment from pGHA2, 100 ng of the EcoRI/Nsil fragment from pTerm2 and 100 ng of a synthetic DNA shown in SEQ ID NO: 15 were dissolved in 20 μl of T4DNA ligase solution, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12 °C for 16 hours. Using the thus prepared recombinant plasmid

DNA, E. coli strain JM109 was transformed to thereby obtain plasmid pMKex1 shown in Fig. 12.

In a separate step, 3 μ g of pAl234 obtained in Fig. 3 was added to 30 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of Pstl was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and dissolved in 20 μ l of T4DNA polymerase I buffer [a buffer containing 33 mM Tris-HCl (pH 8.0), 66 mM potassium acetate, 10 mM magnesium acetate, 0.5 mM DTT and 0.01% BSA]. To the resultant mixture, 5 units of T4DNA polymerase I (Takara Shuzo) was added and reacted at 12 °C for 15 minutes, whereby the 5' cohesive ends generated by the Pstl digestion were changed to blunt ends. The reaction mixture was subjected to phenol-chloroform extraction followed by ethanol precipitation. To the precipitate, 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 100 mM potassium chloride and 1 mM DTT and 10 units of BamHI were added and reacted at 37°C for 4 hours. The reaction mixture was subjected to agarose gel electrophoresis, and about 0.3 μ g of an approx. 0.7 kb DNA fragment containing a cDNA coding for an shIL-5R α fragment was recovered.

Three μg of the expression vector for E. coli, pMKex1 obtained in Fig. 12 was dissolved in 30 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 10 mM magnesium chloride, 100 mM potassium chloride and 1 mM DTT, to which 10 units of BamHl was added and reacted at 37°C for 4 hours. DNA fragments were recovered from the reaction mixture by ethanol precipitation and dissolved in 30 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of EcoRV was added and reacted at 37°C for 4 hours. About 1.5 μg of DNA fragments were recovered from the reaction mixture by ethanol precipitation.

Fifty ng of the thus obtained cDNA coding for an shIL-5R α fragment and 100 ng of the thus obtained RcoRV/BamHI fragment from pMKex1 were dissolved in 20 μ I of T4DNA ligase buffer, to which 200 units of T4DNA ligase was added. Then, ligation was performed at 12°C for 16 hours. Using the thus prepared recombinant plasmid DNA, E. coli strain JM109 was transformed to thereby obtain plasmid pAl263 shown in Fig. 13.

The above plasmid pAl263 was transfected into E. coli (Molecular Cloning, A Laboratory Manual, 2nd Edition published by Cold Spring Harbor Laboratory Press, 1989), which was cultured in 400 ml of LB medium containing 200 μ g/ml of ampicillin at 37°C for 4 hours. Then, 0.5 mM IPTG was added and the cells were cultured at 37°C for another 2 hours. Four hundred ml of the culture medium was centrifuged at 3,000xg for 15 minutes. The precipitate containing the cells of E. coli was suspended in 100 ml of buffer I [10 mM Tris-HCl (pH 8.0), 1 mM EDTA, 150 mM sodium chloride]. After recentrifugation, the precipitate was suspended in 7 ml of buffer I and sonicated to disrupt cells. The resultant suspension was centrifuged at 10,000xg for 30 minutes, and the precipitate was dissolved in 500 μ l of SDS-polyacrylamide gel electrophoresis sample buffer [6 mM Tris-HCl (pH 6.8), 2% SDS, 10% glycerol, 5% 2-mercaptoethanol] and subjected to polyacrylamide gel electrophoresis. Thus, a purified shIL-5R α fragment having a molecular weight of about 27 kD was obtained.

(12) Preparation of a Cell Membrane Fraction from Human IL-5R α Expressing Cells

The preparation of a membrane component from the hIL-5R α gene transfected CTLL-2 cells [J. Exp. Med., <u>177.</u> 1523 (1993)] or control CTLL-2 cells [ATCC TIB 214] was performed as described below.

Briefly, the cells were centrifuged (1,200 rpm, 5 min.), washed with PBS twice, and then suspended in cell disruption buffer [20 mM HEPES (pH 7.4), 1 mM EDTA, 0.5 mM PMSF, 250 mM sucrose] and disrupted with a homogenizer. After the disruption, the cells were centrifuged at 5,500 rpm for 15 minutes to remove the precipitate. The cells were further centrifuged at 35,000 rpm to recover cell membrane fractions as a precipitate.

2. Immunization of Animals and Preparation of Antibody-Producing Cells

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Fifty μ g of each of the antigens obtained in subsections (9), (10), (11) or (12) of section 1 of Example 1 was administered independently to 5-week old female BALB/c mice or female SD rats together with 2 mg of aluminium gel and 1×10^9 cells of pertussis vaccine (Chiba Prefectural Serum Research Institute). 2 weeks after the administration, 50 μ g of the protein was administered once a week in total of 4 times. Blood samples were collected from the venous plexus of eyegrounds or the tail vein, and antibody titer of the serum thereof was examined by the enzyme immunoassay described below under 3. Spleens were removed 3 days after the final immunization from those mice or rats which exhibited a sufficient antibody titer. In this immunization experiment, the cell membrane fraction obtained in subsection (12) of section 1 of Example 1 was used as an antigen to immunize 13 mice and 5 rats. However, no remarkable rise in antibody titer was observed in those animals. Also, no satisfactory rise in antibody titer was observed in the 5 rats immunized with the shIL-5R α obtained in subsection (9) of section 1 of Example 1 or the 10 rats immunized with the shIL-5R α obtained in subsection 1 of Example 1.

The spleen was cut into pieces in MEM medium (Nissui Pharmaceuticals), loosened with tweezers and centrifuged (1,200 rpm, 5 min.). Then, the supernatant was discarded and the remainder was treated with Tris-ammonium chloride buffer (pH 7.65) for 1-2 minutes to remove erythrocytes and washed with MEM medium 3 times. The resultant splenocytes were used for cell fusion.

3. Enzyme Immunoassay

The measurement of antisera or culture supernatants of hybridoma cells derived from mice or rats immunized with the shIL-5R α obtained in subsections (9) or (10) of section 1 of Example 1 was performed according to the two methods described below using, as an antigen, the hIL-5R α -Fc obtained from a culture supernatant of insect cells as described in subsection (10) of section 1 Example 1.

- (A) To a 96-well EIA plate (Greiner), hIL-5R α -Fc diluted to 1 μ g/ml with PBS and a control antigen, anti-GD3 chimeric antibody KM871 having a common human Ig constant region, were dispensed separately in an amount of 50 μ l/well and left at 4°C overnight to have the proteins adsorbed. After washing, PBS containing 1% bovine serum albumin (BSA) (hereinafter, referred to as 1% BSA-PBS) was added to the plate (100 μ l/well) and reacted at room temperature for 1 hour to thereby block the remaining active groups. After discarding 1% BSA-PBS, an immunized mouse or rat-derived antiserum and culture supernatant of a hybridoma were dispensed into the wells (50 μ l/well) and reacted for 2 hours. After washing with Tween-PBS, peroxidase-labeled rabbit anti-mouse immunoglobulin or anti-rat immunoglobulin (DAKO) was added to the plate (50 μ l/well), reacted for 1 hour and washed with Tween-PBS. Thereafter, the resultant mixture was allowed to form a color by using ABTS substrate solution [a solution obtained by dissolving 550 mg of 2,2' azinobis(3-ethylbenzothiazoline-6-sulfonic acid)diammonium salt in 1 L of 0.1 M citrate buffer (pH 4.2) and adding 1 μ l/ml of hydrogen peroxide immediately before use] to measure the absorbance at OD415 nm (NJ2001; Japan Intermed).
- (B) Further, for the purpose of selecting a monoclonal antibody having neutralizing activity against IL-5 with a higher probability, screening was performed for an activity to inhibit binding to an IL-5 receptor by the following procedures using a biotin-labeled human IL-5 and the shIL-5R α -Fc obtained from the insect cell culture supernatant in subsection (10) of section 1 of Example 1. The human IL-5 used for biotin labeling was prepared according to the method described in Journal of Immunological Method, 125, 233 (1989).

The biotin labeling of the human IL-5 was performed according to the protocol attached to a biotin-labeling reagent (Biotin-LC-Hydrazide) (Pierce) by the following procedures. First, 1.6 mg/ml of human IL-5 dissolved in PBS was applied to a PD10 column (Pharmacia) equilibrated with a labeling buffer (100 mM sodium acetate, 0.02% NaN $_3$, pH 5.5) for salt exchange and 1 ml of a fraction having high protein concentration was recovered. To 0.5 ml of this human IL-5 solution, 1 ml of a labeling buffer containing 30 mM metaperiodic acid was added and reacted at room temperature for 30 minutes while shielding the light. After the completion of the reaction, the reaction mixture was applied to a PD10 column equilibrated with a labeling buffer to remove the unreacted metaperiodic acid. Thus, 1.5 ml of a fraction having high protein concentration was recovered. To this fraction, 20 μ l of a labeling buffer containing 5 mM biotin-labeling reagent as described above was added and reacted at room temperature for 1 hour. After the completion of the reaction, 50 μ l of reaction termination buffer (0.1 M Tris, pH 7.5) was added, and then the reaction mixture was applied to a PD10 column equilibrated with 0.05% NaN $_3$ -containing PBS to exchange salts and, simultaneously, remove unreacted reagents. The thus obtained biotin-labeled human IL-5 was stored at 4 °C.

The shIL-5R α -Fc obtained from the insect cell culture supernatant in subsection (10) of section 1 of Example 1 was diluted to a concentration of 5 μ g/ml with PBS, dispensed into a 96-well EIA plate (Greiner) (50 μ l/well) and left at 4 °C overnight to have the protein adsorbed. After washing with PBS, PBS containing 1% bovine serum albumin (BSA) (1% BSA-PBS) was added to the plate (100 μ l/well) and reacted at room temperature for 1 hour to block the remaining active groups. Then, the plate was washed with Tween-PBS. Thereafter, an antiserum derived from immunized mouse or rat and the culture supernatant of the hybridoma, and the biotin-labeled human IL-5 described above were each added to the plate in an amount of 50 μ l/well and reacted at 4 °C overnight. On the next day, the plate was washed with Tween-PBS, and then 50 μ l/well of peroxidase-labeled avidin (Nippon Reizo) diluted 4000 folds with 1% BSA-PBS was added and reacted at room temperature for 1 hour. After washing with Tween-PBS, 50 μ l/well of ABTS substrate solution was added to allow color development and the absorbance at OD415 was measured.

With respect to the measurement of antisera and culture supernatants of hybridomas derived from those mice or rats immunized with the hIL-5R α fragment obtained in subsection (11) of section 1 of Example 1, the hIL-5R α fragment produced by E. coli in subsection (11) of section 1 of Example 1 was used as an antigen. In a manner similar to that described above, the shIL-5R α produced by E. coli and an E. coli cell protein (control antigen) were adsorbed on plates separately. Using thus prepared plates, the reactivity of culture supernatants of hybridomas and antisera of immunized mice or rats was examined.

Further, with respect to the measurement of antisera and culture supernatants of hybridomas derived from those mice or rats immunized with the cell membrane fraction from hIL-5R α expressing cells obtained in subsection (12) of section 1 of Example 1, the cell membrane fraction obtained in subsection (12) of section 1 of Example 1 was used as an antigen. In a manner similar to that described above, the cell membrane fraction from IL-5R α -expressing cells and a cell membrane fraction from control cells were adsorbed on plates separately. Using thus prepared plates, the reactivity of culture supernatants of hybridomas and antisera of immunized mice or rats was examined.

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4. Preparation of Mouse Myeloma Cells

An 8-azaguanine resistant mouse myeloma cell line, P3-U1, was cultured in a normal medium and not less than $2x10^7$ cells were secured and submitted for cell fusion as a parent line.

5. Preparation of Hybridomas

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The mouse or rat splenocytes obtained in section 2 of Example 1 and the myeloma cells obtained in section 4 of Example 1 were mixed at a ratio of 10:1, and the mixture was centrifuged (1,200 rpm, 5 min.). Then, the supernatant was discarded and the precipitated cells were loosened sufficiently. To the resultant cells, a mixed solution composed of 2 g of polyethylene glycol-1000 (PEG-1000), 2 ml of MEM medium and 0.7 ml of DMSO was added in an amount of 0.2 to 1 ml per 10⁸ mouse splenocytes, followed by the addition of 1 to 2 ml portions of MEM medium at 1 to 2 minite interval at 37°C. Thereafter, MEM medium was added to give a total volume of 50 ml. After centrifugation (900 rpm, 5 min.), the supernatant was discarded and cells were loosened gently. Then, cells were gently suspended in 100 ml of HAT medium by suction and release with a pipette.

This cell suspension was dispensed into a 96-well culture plate (100 μ l/well) and cultured in a 5% CO₂ incubator at 37°C for 10-14 days. The resultant culture supernatant was examined by the enzyme immunoassay described in section 3 of Example 1, and those wells which showed specific reaction with the hIL-5R α -Fc prepared from an insect cell culture supernatant or with the shIL-5R α produced by E. coli were selected. Further, the medium was replaced with HT medium and a normal medium, and cloning was repeated twice. As a result, hybridoma cell lines producing an anti-human IL-5R α monoclonal antibody were established.

As a result of screening about 4000 hybridoma clones obtained from 6 mice or 8 rats immunized with the hIL-5R α fragment obtained in subsection (11) of section 1 of Example 1, an anti-human IL-5R α monoclonal antibody was obtained and designated as KM1074. Its reactivity with IL-5R α was extremely weak compared to that of anti-human IL-5R α monoclonal antibodies KM1257 and KM1259 to be described later.

In a separate step, hybridomas were obtained from 12 or 6 animals that exhibited a high antibody titer and which were selected from 15 or 20 mice immunized with the shIL-5R α obtained in subsection (9) of section 1 of Example 1 or the shIL-5R α obtained in subsection (10) of section 1 of Example 1. As a result of screening more than 10000 hybridoma clones, 81 hybridoma clones were established that produced an anti-human IL-5R α monoclonal antibody and which showed a specific reactivity with hIL-5R α expressing cells when tested by the method described later in section 1 of Example 3. Among these, the monoclonal antibody which exhibited the most strong reactivity in the immunocyte staining method described later in section 1 of Example 3 later was KM1257. Hybridoma KM1257 was deposited at the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology (1-3, Higashi 1-Chome, Tsukuba City, Ibaraki, Japan; hereinafter, the address is the same for this Institute) on June 13, 1995 under accession number FERM BP-5133. Of those 81 clones, only six clones exhibited a strong inhibition activity against the biological activity of IL-5 which is described later in section 2 of Example 3. Among these six clones, the monoclonal antibodies which exhibited the strongest inhibition activity were KM1259 and KM1486. Hybridoma KM1259 was deposited under accession number FERM BP-5651 on September 3, 1996 both at the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology.

The reactivities of monoclonal antibodies KM1257, KM1259 and KM1486 are shown in Fig. 14. Subclass of each antibody was determined by an enzyme immunoassay using a subclass typing kit. As a result, the antibody classes of KM1257, KM1259 and KM1486 were all IgG1.

6. Purification of Monoclonal Antibodies

The hybridoma cell line obtained in 5 above was intraperitoneally administered to pristane-treated, female nude mice (Balb/c) of 8 weeks of age at a dose of $(5-20 \times 10^6 \text{ cells/mouse})$. The hybridoma caused ascites tumor 10 to 21 days after the administration. From those mice in which ascites accumulated, ascites was collected (1-8 ml/mouse), centrifuged (3,000 rpm, 5 min.) to remove the solids and then purified by the caprylic acid precipitation method (Antibodies - A Laboratory Manual, Cold Spring Harbor Laboratory, 1988) to obtain purified monoclonal antibody.

EXAMPLE 2

- Preparation of Anti-Human IL-5R α Humanized Antibodies
 - 1. Construction of Tandem Cassette-Type Humanized Antibody Expression Vector pKANTEX93

A tandem cassette-type humanized antibody expression vector, pKANTEX93, for expressing a humanized anti-

body of human antibody IgG1, κ type in animal cells and into which a cDNA coding for a humanized antibody VH and a cDNA coding for a humanized antibody VL were transfected upstream of a cDNA coding for human antibody $C_{\gamma}1$ and a cDNA coding for human antibody C_{κ} , respectively, was constructed as described below based on the plasmid pSE1UK1SEd1-3 disclosed in Kokai No. 257891/90. The humanized antibody expression vector constructed was used for the expression of human chimeric antibodies and human CDR-grafted antibodies in animal cells.

(1) Modification of the Apal and EcoRl Restriction Sites present in Rabbit β-Globin Gene Splicing Signal and Poly (A) Signal

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The modification of the Apal and EcoRI restriction sites present in rabbit β -globin gene splicing poly (A) signal of plasmid pSE1UK1SEd1-3 was performed as described below in order to enable the construction of a human chimeric antibody expression vector or a human CDR-grafted antibody (=humanized antibody) expression vector by inserting into a humanized antibody expression vector the variable region of a human chimeric antibody or a human CDR-grafted antibody in a cassette using a Notl-Apal fragment (VH) and an EcoRI-Sp1I fragment (VL).

Briefly, 3 μ g of plasmid pBluescript SK(-) (Stratagene) was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the 3' sticky ends generated by the Apal digestion were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSA shown in Fig. 15.

Further, 3 μ g of the thus obtained plasmid pBSA was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the 5' sticky ends generated by the EcoRl digestion were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using the thus obtained recombinant plasmid DNA solution, \underline{E} , \underline{coli} HB101 was transformed to obtain plasmid pBSAE shown in Fig. 16.

Subsequently, 3 μ g of the thus obtained plasmid pBSAE was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT, to which 10 units of the restriction enzyme HindIII (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 20 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT. The resultant mixture was divided into two 10 μ l portions. To one portion, 10 units of the restriction enzyme SacII (Toyobo) was added, and to the other portion, 10 units of the restriction enzyme KpnI (Takara Shuzo) was added. Then, both mixtures were reacted at 37 °C for 1 hour. Both reaction mixtures were subjected to agarose gel electrophoresis, and an approx. 2.96 kb HindIII-SacII fragment and an approx. 2.96 kb KpnI-HindIII fragment were recovered, each in about 0.3 μ g.

Subsequently, 3 μ g of plasmid pSE1UK1SEd1-3 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme SacII (Toyobo) and 10 units of the restriction enzyme KpnI (Takara Shuzo) were added and reacted at 37 °C for 1 hour. The reaction mixture was eth-anol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 50 mM sodium chloride and 1 mM DTT. To the resultant mixture, 10 units of the restriction enzyme HindIII (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis, and an approx. 2.42 kb HindIII-SacII fragment and an approx. 1.98 kb KpnI-HindIII fragment were recovered, each in about 0.2 μ g.

Then, 0.1 μ g of the HindIII-SacII fragment from plasmid pSE1UK1SEd1-3 and 0.1 μ g of the HindIII-SacII fragment from pBSAE obtained above were dissolved in sterilized water to give a total volume of 20 μ I and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSH-S shown in Fig. 17. Also, 0.1 μ g of the KpnI-HindIII fragment from plasmid pSE1UK1SEd1-3 and 0.1 μ g of the KpnI-HindIII fragment from pBSAE obtained above were dissolved in sterilized water to give a total volume of 20 μ I and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSK-H shown in Fig. 18.

Subsequently, 3 μ g each of the thus obtained plasmids pBSH-S and pBSK-H were added separately to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. Both reaction mixtures were ethanol-precipitated, and the 3' sticky ends generated by the Apal digestion were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using each of the thus obtained recombinant plasmid DNA solutions, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSH-SA and pBSK-HA shown in Fig. 19.

Subsequently, 5 μg each of the thus obtained plasmids pBSH-SA and pBSK-HA were added separately to 10 μl of

a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37°C for 10 minutes so that the plasmid was partially digested. Then, both reaction mixtures were ethanol-precipitated. After the 5' sticky ends generated by the EcoRl digestion were blunted using DNA Blunting Kit (Takara Shuzo), both reaction mixtures were subjected to agarose gel electrophoresis, and an approx. 5.38 kb fragment and an approx. 4.94 kb fragment were recovered, each in about 0.5 μ g. Then, 0.1 μ g each of the thus recovered fragments were dissolved separately in sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using each of the thus obtained recombinant plasmid DNA solutions, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmids pBSH-SAE and pBSK-HAE shown in Fig. 20.

Subsequently, 3 μ g each of the thus obtained plasmids pBSH-SAE and pBSK-HAE were added separately to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride, 100 mM sodium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. Both reaction mixtures were ethanol-precipitated and the 5' sticky ends generated by the EcoRl digestion were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using each of the thus obtained recombinant plasmid DNA solutions, <u>E. coli</u> HB101 was transformed to obtain plasmids pBSH-SAEE and pBSK-HAEE shown in Fig. 21. Ten μ g each of the thus obtained plasmids were separately reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that both the Apal and EcoRl restriction sites had been eliminated by the above-described modification.

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(2) Introduction of a Sall Restriction Site into the Downstream Portion consisting of the Rabbit β-Globin Gene Splicing Signal, Rabbit β-Globin Gene Poly (A) Signal and SV40 Early Gene Poly (A) Signal

In order to ensure that expression promoters for the human antibody H and L chains in a humanized antibody expression vector could be replace with any promoters, a Sall restriction site was transfected into the downstream portion consisting of the rabbit β -globin gene splicing signal, rabbit β -globin gene poly (A) signal and SV40 early gene poly (A) signal of plasmid pSE1UK1SEd1-3 as described below.

Briefly, 3 μg of the plasmid pBSK-HAEE obtained in subsection (1) of section 1 of Example 2 was added to 10 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Nael (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was dissolved in 20 μl of a buffer containing 50 mM Tris-HCl (pH 9.0) and 1 mM magnesium chloride, to which 1 unit of alkaline phosphatase (E. coli C75, Takara Shuzo) was added and reacted at 37 °C for 1 hour to dephosphorylate 5' ends. Then, the reaction mixture was subjected to phenol-chloroform extraction, followed by ethanol precipitation. The precipitate was dissolved in 20 μl of a buffer containing 10 mM Tris-HCl (pH 8.0) and 1 mM ethylenediamine-tetraacetic acid disodium (hereinafter referred to as "TE buffer"). One μl of the mixture and 0.1 μg of a phosphorylated Sall linker (Takara Shuzo) were added to sterilized water to give a total volume of 20 μl, and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to obtain plasmids pBSK-HAEESal shown in Fig. 22. Ten μg each of the thus obtained plasmid was reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequence (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that one Sall restriction site had been transfected into the downstream portion consisting of the rabbit β -globin gene splicing signal, rabbit β -globin gene poly (A) signal and SV40 early gene poly (A) signal.

(3) Modification of the Apal Restriction Site present in the Poly (A) Signal of Herpes Simplex Virus Thymidine Kinase (hereinafter referred to as "HSVtk") Gene

The modification of the Apal restriction site present in the poly (A) signal of HSVtk gene located downstream of Tn5 kanamycin phosphotransferase gene in plasmid pSE1UK1SEd1-3 was performed as described blow.

Briefly, 3 μ g of the plasmid pBSA obtained in subsection (1) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme SacIl (Toyobo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis and about 1 μ g of an approx. 2.96 kb SacIl-Xhol fragment was recovered.

Subsequently, $5~\mu g$ of plasmid pSE1UK1SEd1-3 was added to 10 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme SacII (Toyobo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was dissolved in 10 μl of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to

which 10 units of the restriction enzyme XhoI (Takara Shuzo) was added and reacted at 37° C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis and about 1 μ g of an approx. 4.25 kb SacII-XhoI fragment was recovered.

Subsequently, 0.1 μg of the SacII-XhoI fragment from pBSA and the SacII-XhoI fragment from plasmid pSE1UK1SEd1-3 as obtained above were added to sterilized water to give a total volume of 20 μ I, and then ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSX-S shown in Fig. 23.

Subsequently, 3 μ g of the thus obtained plasmid pBSX-X was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the 3' sticky ends generated by the Apal digestion were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSX-SA shown in Fig. 24. Ten μ g of the thus obtained plasmid was reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that the Apal restriction site of the HSVtk gene poly (A) signal had been eliminated.

(4) Construction of a Humanized Antibody L Chain Expression Unit

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Plasmid $mMohC\kappa$ having a humanized antibody L chain expression unit in which a cDNA coding for the constant region of human κ -type L-chain ($C\kappa$) was located downstream of the promoter/enhancer of the terminal repeated sequence of Moloney mouse leukemia virus and into which a cDNA coding for VL of a human chimeric antibody or human CDR-grafted antibody could be inserted in a cassette was constructed as described below.

Briefly, 3 μg of plasmid pBluescript SK(-) (Stratagene) was added to 10 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Sacl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Clal (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the sticky ends generated by the Sacl and Clal digestions were blunted using DNA Blunting Kit (Takara Shuzo). Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μg of an approx. 2.96 kb DNA fragment. Then, 0.1 μg of the recovered DNA fragment was added to sterilized water to give a total volume of 20 μl and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . coli HB101 was transformed to obtain plasmid pBSSC shown in Fig. 25.

Subsequently, 3 μ g of the thus obtained plasmid pBSSC was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 2.96 kb Kpnl-Xhol fragment.

Subsequently, 5 μ g of the plasmid pAGE147 disclosed in Kokai No. 205694/94 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme KpnI (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.3 μ g of an approx. 0.66 kb KpnI-Xhol fragment containing the promoter/enhancer of the terminal repeated sequence of Moloney mouse leukemia virus.

Subsequently, 0.1 μ g of the Kpnl-Xhol fragment from pBSSC and 0.1 μ g of the Kpnl-Xhol fragment from pAGE147 as obtained above were dissolved in sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBSMo shown in Fig. 26.

Subsequently, 3 μ g of the plasmid pBSMo obtained above was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added further and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme HindIII (Takara Shuzo) was added further and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover

about 1 μg of an approx. 3.62 kb Kpnl-HindIII fragment.

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Subsequently, two synthetic DNAs having the base sequences shown in SEQ ID NOS: 16 and 17, respectively, were synthesized using an automatic DNA synthesizer (380A, Applied Biosystems). Then, 0.3 μ g each of the synthetic DNAs obtained were added to 15 μ l of sterilized water and heated at 65°C for 5 minutes. The reaction mixture was left at room temperature for 30 minutes. To this mixture, 2 μ l of a 10x buffer [500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT] and 2 μ l of 10 mM ATP were added. Further, 10 units of T4 polynucleotide kinase (Takara Shuzo) was added and reacted at 37°C for 30 minutes to phosphorylate the 5' ends. Then, 0.1 μ g of the Kpnl-HindIII fragment (3.66 kb) from plasmid pBSMo as obtained above and 0.05 μ g of the phosphorylated synthetic DNAs were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, μ c. coli HB101 was transformed to obtain plasmid pBSMoS shown in Fig. 27. Ten μ g of the thus obtained plasmid was reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that the synthetic DNAs of interest had been transfected.

Subsequently, 3 μ g of the plasmid pChilgLA1 disclosed in Kokai No. 304989/93 was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units each of the restriction enzymes EcoRl (Takara Shuzo) and RcoRV (Takara Shuzo) were added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 9.70 kb EcoRl-EcoRV fragment. Subsequently, two synthetic DNAs having the base sequences shown in SEQ ID NOS: 18 and 19, respectively, were synthesized using an automatic DNA synthesizer (380A, Applied Biosystems). Then, 0.3 μ g each of the obtained synthetic DNAs were added to 15 μ l of sterilized water and heated at 65°C for 5 minutes. The reaction mixture was left at room temperature for 30 minutes. To this solution, 2 μ l of a 10x buffer [500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT] and 2 μ l of 10 mM ATP were added. Further, 10 units of T4 polynucleotide kinase (Takara Shuzo) was added and reacted at 37°C for 30 minutes to phosphorylate the 5' ends. Then, 0.1 μ g of the EcoRl-EcoRV fragment (9.70 kb) from plasmid pChilgLA1 as obtained above and 0.05 μ g of the phosphorylated synthetic DNAs were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . coli HB101 was transformed to obtain plasmid pChilgLA1S shown in Fig. 28.

Subsequently, 3 μ g of the plasmid pBSMoS as obtained above was dissolved in 10 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 100 mM potassium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Hpal (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to recover about 1 μ g of an approx. 3.66 kb Hpal-EcoRl fragment.

Subsequently, 10 μ g of the plasmid pChilgLA1S as obtained above was dissolved in 10 μ l of a buffer containing 20 mM Tris-HCl (pH 7.9), 50 mM potassium acetate, 10 mM magnesium acetate, 1 mM DTT and 100 μ g/ml BSA, to which 10 units of the restriction enzyme NlaIV (New England Biolabs) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to recover about 0.3 μ g of an approx. 0.41 kb NlaIV-EcoRl fragment.

Subsequently, 0.1 μg each of the Hpal-EcoRI fragment from pBSMoS and the NIaIV-EcoRI fragment from pChilgLA1S as obtained above were added to sterilized water to give a total volume of 20 μ I and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pMohC κ shown in Fig. 29.

(5) Construction of a Humanized Antibody H Chain Expression Unit

Plasmid $mMohC_{\gamma}$ 1 having a humanized antibody H chain expression unit in which a cDNA coding for the constant region of human lgG1 type H-chain ($C_{\gamma}1$) was located downstream of the promoter/enhancer of the terminal repeated sequence of Moloney mouse leukemia virus and into which a cDNA coding for VH of a human chimeric antibody or human CDR-grafted antibody could be inserted in a cassette was constructed as described below.

Briefly, 3 μg of the plasmid pBSMo obtained in subsection (4) of section 1 of Example 2 was added to 10 μl of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μl of a buffer containing 30 mM sodium acetate (pH 5.0), 100 mM sodium chlorids, 1 mM zinc acetate and 10% glycerol, to which 10 units of mung bean nuclease (Takara Shuzo) was added and reacted at 37°C for 10 minutes. The reaction mixture was subjected to phenol-chloro-

form extraction, followed by ethanol precipitation. Then, the sticky ends were blunted using DNA Blunting Kit (Takara Shuzo) and the resultant DNA fragments were ligated using DNA Ligation Kit (Takara Shuzo). Using the thus obtained recombinant plasmid DNA solution, <u>E. coli</u> HB101 was transformed to obtain plasmid pBSMoSal shown in Fig. 30. Ten µg of the thus obtained plasmid was reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that the Xhol restriction site located upstream of the promoter/enhancer of the terminal repeated sequence of Moloney mouse leukemia virus had been eliminated.

Subsequently, 3 μ g of the plasmid pBSMosal as obtained above was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme HindIII (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 3.66 kb Kpnl-HindIII fragment.

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Subsequently, two synthetic DNAs having the base sequences shown in SEQ ID NOS: 20 and 21, respectively, were synthesized using an automatic DNA synthesizer (380A, Applied Biosystems). Then, 0.3 μ g each of the obtained synthetic DNAs were added to 15 μ l of sterilized water and heated at 65°C for 5 minutes. The reaction mixture was left at room temperature for 30 minutes. To this solution, 2 μ l of a 10x buffer [500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT] and 2 μ l of 10 mM ATP were added. Further, 10 units of T4 polynucleotide kinase (Takara Shuzo) was added and reacted at 37°C for 30 minutes to phosphorylate the 5' ends. Then, 0.1 μ g of the Kpnl-HindIII fragment (3.66 kb) from plasmid pBSMoSal as obtained above and 0.05 μ g of the phosphorylated synthetic DNAs were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, μ c. coli HB101 was transformed to obtain plasmid pBSMoSalS shown in Fig. 31. Ten μ g of the thus obtained plasmid was reacted according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech) to thereby determine the base sequence. As a result, it was confirmed that the synthetic DNAs of interest had been transfected.

Subsequently, 10 μ g of the plasmid pChilgHB2 disclosed in Kokai No. 304989/93 was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Eco52l (Toyobo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 30 mM sodium acetate (pH 5.0), 100 mM sodium chloride, 1 mM zinc acetate and 10% glycerol, to which 10 units of mung bean nuclease (Takara Shuzo) was added and reacted at 37°C for 10 minutes. The reaction mixture was subjected to phenol-chloroform extraction, followed by ethanol precipitation. Then, the sticky ends were blunted using DNA Blunting Kit (Takara Shuzo). After ethanol precipitation, the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.7 μ g of an approx. 0.99 kb Apal-blunt end fragment.

Subsequently, 3 μ g of plasmid pBluescript SK(-) (Stratagene) was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 33 mM Tris-HCl (pH 7.9), 10 mM magnesium acetate, 66 mM potassium acetate, 0.5 mM DTT and 10 μ g/ml BSA, to which 10 units of the restriction enzyme Smal (Takara Shuzo) was added and reacted at 30 °C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 3.0 kb Apal-Smal fragment.

Subsequently, 0.1 μ g of the Apal-blunt end fragment from plasmid pChilgHB2 as obtained above and 0.1 μ g of the Apal-Smal fragment from pBluescript SK(-) were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pBShC γ 1 shown in Fig. 32.

Subsequently, 5 μ g of the plasmid pBShC γ 1 as obtained above was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Spel (Takara Shuzo) was added and reacted at 37 °C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 1.0 kb Apal-Spel fragment.

Subsequently, 3 μ g of the plasmid pBSMoSalS as obtained above was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate

was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Spel (Takara Shuzo) was added and reacted at 37 °C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 3.66 kb Apal-Spel fragment.

Subsequently, 0.1 μ g each of the Apal-Spel fragment from pBShC γ 1 and the Apal-Spel fragment from pBSMoSalS as obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to obtain plasmid pMohC γ 1 shown in Fig. 33.

(6) Construction of Tandem Cassette-Type Humanized Antibody Expression Vector pKANTEX93

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A tandem cassette-type humanized antibody expression vector, pKANTEX93, was constructed as follows using the various plasmids obtained in subsections (1)-(5) of section 1 of Example 2.

Briefly, 3 μ g of the plasmid pBSH-SAEE obtained in subsection (1) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme HindIII (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Sall (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 5.42 kb HindIII-Sall fragment.

Subsequently, 5 μ g of the plasmid pBSK-HAEE obtained in subsection (1) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme HindIII (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.8 μ g of an approx. 1.98 kb Kpnl-HindIII fragment containing rabbit β -globin gene splicing poly (A) signal, SV40 early gene poly (A) signal and SV40 early gene promoter.

Subsequently, 5 μ g of the plasmid pMohC γ 1 obtained in subsection (5) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Sall (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.8 μ g of an approx. 1.66 kb Kpnl-Sall fragment containing the humanized antibody H chain expression unit.

Subsequently, 0.1 μ g each of the HindIII-Sall fragment from pBSH-SAEE, the KpnI-HindIII fragment from pBSK-HAEE and the KpnI-SalI fragment from pMohC γ 1 as obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pMo γ 1SP shown in Fig. 34.

Subsequently, 3 μ g of the thus obtained plasmid pMo γ 1SP was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units each of the restriction enzymes Sall (Takara Shuzo) and Xhol were added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 9.06 kb Sall-Xhol fragment.

Subsequently, 5 μ g of the plasmid pBSK-HAEESal obtained in subsection (2) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was eth-anol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Sall (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electro-phoresis to thereby recover about 0.7 μ g of an approx. 1.37 kb Kpnl-Sall fragment containing rabbit β -globin gene splicing signal, rabbit β -globin gene splicing signal poly (A) signal and SV40 early gene poly (A) signal.

Subsequently, 5 μ g of the plasmid pMohC κ obtained in subsection (4) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Kpnl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.7 μ g of an approx. 1.06 kb Kpnl-Xhol fragment containing the humanized antibody L chain

expression unit.

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Subsequently, 0.1 μ g each of the Sall-Xhol fragment from pMo γ 1SP, the Kpnl-Sall fragment from pBSK-HAEESal and the Kpnl-Xhol fragment from pMohC κ as obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to obtain plasmid pMo $\kappa\gamma$ 1SP shown in Fig. 35.

Subsequently, 3 μg of the thus obtained plasmid pMo $\kappa\gamma$ 1SP was dissolved in 10 μl of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme SacII (Toyobo) was added and reacted at 37 °C for 10 minutes so that the DNA fragments were partially digested. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.2 μg of an approx. 8.49 kb SacII-Xhol fragment.

Subsequently, 3 μ g of plasmid pBSX-SA obtained in subsection (3) of section 1 of Example 2 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme SacII (Toyobo) was added and reacted at 37 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was dissolved in 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Xhol (Takara Shuzo) was added and reacted at 37°C for 1 hour. Then, the reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 4.25 kb SacII-Xhol fragment.

Subsequently, 0.1 μ g each of the SacIl-Xhol fragment from pMo $\kappa\gamma$ 1SP and the SacIl-Xhol fragment from pBSX-SA as obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pKANTEX93 shown in Fig. 36.

- 2. Isolation and Analysis of the cDNAs Coding for Anti-Human IL-5R α Monoclonal Antibodies
 - (1) Isolation of mRNA from Anti-Human IL-5R α Monoclonal Antibody-Producing Hybridomas

Using Fast Track, an mRNA extraction kit manufactured by Invitrogen, mRNA was isolated from $1x10^8$ cells each of mouse anti-human IL-5R α monoclonal antibodies KM1257, KM1259 and KM1486 producing hybridoma cell lines (corresponding to hybridomas FERM BP-5133, FERM BP-5134 and FERM BP-5651, respectively) in accordance with the instructions attached to the kit.

(2) Preparation of H and L Chain cDNA Libraries from Mouse Anti-Human IL-5R α Monoclonal Antibody-Producing Hybridomas

Using cDNA Synthesis Kit (Pharmacia Biotech) and according to the instructions attached to the kit, a cDNA having an EcoRI adapter at both ends was synthesized separately from 5 μ g each of the mRNAs obtained from KM1257, KM1259 and KM1486 in subsection (1) of section 2 of Example 2. About 6 μ g of each cDNA was dissolved in 10 μ l of sterilized water and subjected to agarose gel electrophoresis, to thereby recover about 0.1 μ g each of an approx. 1.5 kb cDNA fragment corresponding to the cDNA encoding for the H chain of IgG type antibody and an approx. 1.0 kb fragment corresponding to the L chain of immunoglobulins. Then, 0.1 μ g of the approx. 1.5 kb cDNA fragment or the approx. 1.0 kb cDNA fragment and 1 μ g of Lamda ZAPII vector [as treated with calf intestine alkaline phosphatase after cleavage with EcoRI; Stratagene) were dissolved in 11.5 μ l of T4 ligase buffer, to which 175 units of T4 DNA ligase was added and incubated at 12°C for 24 hours, followed by incubation at room temperature for another 2 hours. Using 4 μ l of each reaction mixture, cDNAs were packed into a λ phage using Giga Pack Gold (Stratagene) by conventional methods (Molecular Cloning, 2.95, Cold Spring Harbor Laboratory, 1989). The resultant λ phages were infected to E. coli strain XL1-Blue [Biotechniques, E, 376 (1987)] in Giga Pack Gold by conventional methods (Molecular Cloning, 2.95-107, Cold Spring Harbor Laboratory, 1989) to obtain about 4000 phage clones for each of the H chain cDNA library and the L chain cDNA library of KM1257, KM1259 and KM1486.

(3) Cloning of the cDNAs Coding for the H and L Chains of Anti-Human IL-5R α Monoclonal Antibody-Producing Hybridomas

The recombinant phages prepared in subsection (2) of section 2 of Example 2 was fixed on a nitrocellulose filter by conventional methods (Molecular Cloning, 2.12, Cold Spring Harbor Laboratory, 1989). The cDNA coding for the C region of mouse Ig {the H chain was a fragment from mouse $C\gamma$ 1 cDNA [Cell, 18, 559 (1979)] and the L chain was a fragment from mouse $C\kappa$ cDNA [Cell, 22, 197 (1980)]} were labeled using ECL direct nucleic acid labelling and detection systems (Amersham). Using those labeled cDNA as probes, recombinant phages were screened. Subsequently,

according to the instructions attached to Lamda ZAPII vector (Stratagene), the phage clones were replaced with plasmid pBluescriptSK(-). Finally, the following plasmids were obtained: recombinant plasmid pKM1257H comprising a cDNA coding for the H chain of KM1257 and recombinant plasmid pKM1257L comprising a cDNA coding for the L chain of KM1259; recombinant plasmid pKM1259H comprising a cDNA coding for the H chain of KM1259 and recombinant plasmid pKM1259L comprising a cDNA coding for the L chain of KM1259; and recombinant plasmid pKM1486H comprising a cDNA coding for the H chain of KM1486 and recombinant plasmid pKM1486L comprising a cDNA coding for the L chain of KM1486.

(4) Determination of the Base Sequences for the V Regions of the cDNAs Coding for the H and L Chains of Anti-Human IL-5R α Monoclonal Antibodies

The base sequence for the V region of each of the cDNAs coding for the H and L chains of mouse anti-human IL-5R α monoclonal antibodies as obtained in subsection (3) of section 2 of Example 2 was analyzed by reacting 10 μ g of the resultant plasmid according to the recipe attached to AutoRead Sequencing Kit (Pharmacia Biotech) and then electrophoresed with A.L.F. DNA Sequencer (Pharmacia Biotech). From the base sequence thus determined for each of the cDNAs, amino acid sequences for the V regions of the L and H chains of KM1257, KM1259 and KM1486 were determined. SEQ ID NO: 22 shows the base sequence and amino acid sequence of the V region of the H chain of KM1257; SEQ ID NO: 23 shows those of the L chain of KM1257; SEQ ID NO: 24 shows those of the H chain of KM1259; SEQ ID NO: 25 shows those of the L chain of KM1259; SEQ ID NO: 26 shows those of the H chain of KM1486; and SEQ ID NO: 27 shows those of the L chain of KM1486.

(5) Identification of CDR sequences for the H and L Chains of Anti-Human IL-5R α Monoclonal Antibodies

CDR sequence for each H chain and those for each L chain were identified from the amino acid sequences for the V regions of the H and L chains of each mouse anti-human IL-5R α as monoclonal antibody determined in subsection (4) of section 2 of Example 2 by comparing the above amino acid sequences with the V region amino acid sequences for known antibodies (Sequences of Proteins of Immunological Interest, US Dept. Health and Human Services, 1991). SEQ ID NOS: 28, 29 and 30 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the H chain of KM1257. SEQ ID NOS: 31, 32 and 33 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the L chain of KM1257. SEQ ID NOS: 34, 35 and 36 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the H chain of KM1259. SEQ ID NOS: 37, 38 and 39 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the L chain of KM1259. SEQ ID NOS: 40, 41 and 42 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the H chain of KM1486. SEQ ID NOS: 43, 44 and 45 show the amino acid sequences for CDR1, CDR2 and CDR3, respectively, of the L chain of KM1486.

3. Preparation of Anti-Human IL-5R α Human Chimeric Antibody

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An anti-human IL-5R α human chimeric antibody derived from the anti-human IL-5R α monoclonal antibody KM1259 having an activity to inhibit the biological activity of human IL-5 was prepared as described below.

(1) Construction of Expression Vector pKANTEX1259 for Anti-Human IL-5R α Human Chimeric Antibody

An expression vector, pKANTEX1259, for an anti-human IL-5R α human chimeric antibody was constructed as follows using the humanized antibody expression vector pKANTEX93 constructed in section 1 of Example 2 and the plasmids pKM1259H and pKM1259L obtained in section 2 of Example 2.

Briefly, 3 μ g of the humanized antibody expression vector pKANTEX93 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT, 100 μ g/ml BSA and 0.01% Triton X-100, to which 10 units of the restriction enzyme Notl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 12.75 kb Apal-Notl fragment. Subsequently, 5 μ g of plasmid pKM1259H was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Banl (Toyobo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT, 100 μ g/ml BSA and 0.01% Triton X-100, to which 10 units of the restriction enzyme Notl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.5 μ g of an approx. 0.41 kb Banl-Notl fragment.

Subsequently, two synthetic DNAs having the base sequences shown in SEQ ID NOS: 46 and 47, respectively,

were synthesized with an automatic DNA synthesizer (380A, Applied Biosystems). Then, 0.3 μg each of the obtained synthetic DNAs were added to 15 μl of sterilized water and heated at 65°C for 5 minutes. After the reaction mixture was left at room temperature for 30 minutes, 2 μl of a 10x buffer [500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT] and 2 μl of 10 mM ATP were added. Further, 10 units of T4 polynucleotide kinase was added and reacted at 37 °C for 30 minutes to thereby phosphorylate the 5' ends.

Then, 0.1 μg of the Apal-Notl fragment from the humanized antibody expression vector pKANTEX93, 0.1 μg of the Banl-Notl fragment from plasmid pKM1259H and 0.05 μg of the phosphorylated synthetic DNAs as obtained above were added to sterilized water to give a total volume of 20 μl and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pKANTEX1259H shown in Fig. 37.

Subsequently, 3 μ g of the thus obtained plasmid pKANTEX1259H was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5) 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT and 100 μ g/ml BSA, to which 10 units each of the restriction enzymes EcoRl (Takara Shuzo) and Spll (Takara Shuzo) were added and reacted at 37 °C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 13.20 kb EcoRl-Spll fragment.

Subsequently, 5 μ g of plasmid pKM1259L was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Avall (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme EcoRl (Takara Shuzo) was added and reacted at 37 °C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.5 μ g of an approx. 0.38 kb Avall-EcoRl fragment.

Subsequently, two synthetic DNAs having the base sequences shown in SEQ ID NOS: 48 and 49, respectively, were synthesized with an automatic DNA synthesizer (380A, Applied Biosystems). Then, 0.3 μ g each of the obtained synthetic DNAs were added to 15 μ l of sterilized water and heated at 65°C for 5 minutes. After the reaction mixture was left at room temperature for 30 minutes, 2 μ l of a 10x buffer [500 mM Tris-HCl (pH 7.6), 100 mM magnesium chloride, 50 mM DTT] and 2 μ l of 10 mM ATP were added. Further, 10 units of T4 polynucleotide kinase was added and reacted at 37 °C for 30 minutes to thereby phosphorylate the 5' ends.

Then, 0.1 μg of the EcoRI-SpII fragment from plasmid KANTEX1259H, 0.1 μg of the AvaII-EcoRI fragment from plasmid pKM1259L and 0.05 μg of the phosphorylated synthetic DNAs as obtained above were added to sterilized water to give a total volume of 20 μl and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, \underline{E} . \underline{coli} HB101 was transformed to obtain plasmid pKANTEX1259 shown in Fig. 38.

(2) Expression of Anti-Human IL-5 α Human Chimeric Antibody in Rat Myeloma YB2/0 Cells (ATCC CRL1581) using pKANTEX1259

The transfection of the anti-human IL-5 α human chimeric antibody expression vector pKANTEX1259 into YB2/0 cells was performed according to the method of Miyaji et al. by electroporation [Cytotechnology,3, 133, (1990)].

Briefly, 4 μg of the pKANTEX1259 obtained in subsection (1) of section 3 of Example 2 was transfected into 4x10⁶ YB2/0 cells. Then, RPMI1640-FCS(10) was dispensed into a 96-well microtiter plate (200 μ l/well). Cells were cultured in a 5% CO₂ incubator at 37°C for 24 hours. Then, Geneticin (hereinafter referred to as "G418"; Gibco) was added to give a concentration of 0.5 mg/ml and cells were cultured for another 1-2 weeks. The culture supernatants were recovered from those wells which had become confluent with the appearance of transformant colonies having G418 resistance. The activity of an anti-human IL-5R α human chimeric antibody in the supernatants was determined by ELISA method 1 or 2 as described below.

ELISA method 1

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The shIL-5R α -Fc obtained from the insect cell culture supernatant in subsection (10) of section 1 of Example 1 was diluted with PBS to a concentration of 5 μ g/ml or less. The diluent was dispensed into a 96-well EIA plate (Greiner) (50 μ l/well), which was left at 4°C overnight to allow the protein to be adsorbed. After washing the plate, PBS containing 1% bovine serum albumin (BSA)(1% BSA-PBS) was added to the plate in an amount of 100 μ l/well and reacted at room temperature for 1 hour to thereby block the remaining active groups. After discarding 1% BSA-PBS, the culture supernatants from the transformant or various purified anti-human IL-5 α antibodies at a concentration of 40 μ g/ml were added to the plate in an amount of 25 μ l/well. Further, the biotin-labeled human IL-5 (0.4 μ g/ml) prepared in section 3 of Example 1 was added to the plate in an amount of 25 μ l/well and reacted at room temperature for 4 hours. After washing with 0.05% Tween-PBS, peroxidase-labeled avidin D (Nippon Reizo) diluted 4000 folds with 1% BSA-PBS was added to the plate in an amount of 50 μ l/well and reacted at room temperature for 1 hour. After washing with 0.05%

Tween-PBS, an ABTS substrate solution [as prepared by dissolving 550 mg of 2,2' azinobis(3-ethylbenzothiazo line-6-sulfonic acid)diammonium in 1 L of 0.1 M citrate buffer (pH 4.2) and adding 1 μ l/ml of hydrogen peroxide immediately before use] was added at 50 μ l/well to allow color development. Then, the absorbance (OD) at 415 nm was measured. The absorbance value in the absence of an antibody was regarded as zero percent inhibition, and the percent inhibitions of antibodies against the biotin-labeled IL-5 were calculated by the following formula to evaluate each sample.

Percent binding inhibition =
$$100 - \frac{A - C}{B - C} \times 100$$

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A: OD value in the presence of an antibody

B: OD value in the absence of an antibody

C: OD value in the absence of biotin-labeled human IL-5.

ELISA method 2

The shIL-5R α obtained from the insect cell culture supernatant in subsection (10) of section 1 of Example 1 was diluted with PBS to a concentration of 2 μ g/ml or less. The diluent was dispensed into a 96-well EIA plate (Greiner) (50 μ l/well), which was left at 4°C overnight to allow the protein to be adsorbed. After washing the plate, PBS containing 1% bovine serum albumin (BSA) (1% BSA-PBS) was added to the plate in an amount of 100 μ l/well and reacted at room temperature for 1 hour to thereby block the remaining active groups. After discarding 1% BSA-PBS, the culture supernatants from the transformant or various purified anti-human IL-5 α antibodies at a concentration of 50 μ g/ml were added to the plate in an amount of 50 μ l/well and reacted at room temperature for 2 hours. After washing with 0.05% Tween-PBS, peroxidase-labeled anti-human IgG antibody (American Qualex International, Inc.) diluted 500 folds with 1% BSA-PBS was added to the plate in an amount of 50 μ l/well and reacted at room temperature for 1 hour. After washing with 0.05% Tween-PBS, an ABTS substrate solution [as prepared by dissolving 550 mg of 2,2' azinobis(3-ethylbenzothiazoline-6-sulfonic acid)diammonium in 1 L of 0.1 M citrate buffer (pH 4.2) and adding 1 μ l/ml of hydrogen peroxide immediately before use] was added at 50 μ l/well to allow color development. Then, the absorbance (OD) at 415 nm was measured.

Those transformant in which the activity of anti-human IL-5R α human chimeric antibody was observed in their culture supernatants were suspended in RPMI1640-FCS(10) medium containing 0.5 mg/ml G418 and 50 nM MTX (Sigma), and cultured in a 5% CO_2 incubator at 37 °C for 1-2 weeks, to thereby induce transformant having resistance to 50 nM MTX. When transformant became confluent in wells, the activity of anti-human IL-5R α human chimeric antibody in the supernatant was measured by either of the ELISA methods described above. Those transformant in which the activity was observed were further cultured in a manner similar to that described above, with the MTX concentration increased to 100 nM and to 200 nM. Thus, transformant which could grow in RPMI1640-FCS(10) medium containing 0.5 mg/ml G418 and 200 nM MTX and which produced an anti-human IL-5R α human chimeric antibody were obtained. The thus obtained transformant were subjected to cloning by the applications of the limiting dilution method to thereby obtain final anti-human IL-5R α human chimeric antibody-producing transformant. As a specific example of the antihuman IL-5R α human chimeric antibody produced by this strain was designated as KM1399. The transformant KM1399 was deposited with the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on September 3, 1996 under accession number FERM BP-5650. The productivity of the anti-human IL-5R α human chimeric antibody KM1399 in the transformant clone KM1399 was approximately 5 μ g/10⁶ cells/24 hr.

(3) Purification of the Anti-Human IL-5R α Human Chimeric Antibody KM1399 from Culture Supernatant

The anti-human IL-5R α human chimeric antibody KM1399 obtained in subsection (2) of section 3 of Example 2 was suspended in GIT medium (Nippon Pharmaceuticals) containing 0.5 mg/ml G418 and 200 mM MTX to give a concentration of 1-2 x 10^5 cells/ml, and dispensed in 200 ml portions into 175 cm² flasks (Greiner). The cells were cultured in a 5% CO₂ incubator at 37°C for 5-7 days, and the culture supernatant was recovered when each flask became confluent. From about 1.0 liter of the culture supernatant, about 3 mg of purified anti-human IL-5R α human chimeric antibody KM1399 was obtained using a Procep A (Bioprocessing) column. About 4 μ g of the purified anti-human IL-5R α human chimeric antibody KM1399 was electrophoresed according to known methods [Nature, 227, 680 (1970)] to perform molecular weight analyses. The results are shown in Fig. 39. As seen from Fig. 39, the molecular weight of the antibody H chain was about 50 KDa and that of the antibody L chain about 25 KDa under reducing conditions. Thus, the expression of the H and L chains with correct molecular weights was confirmed. On the other hand, under non-

reducing conditions, the molecular weight of the anti-human IL-5R α human chimeric antibody KM1399 was about 140 KDa. Thus, the expression of a human chimeric antibody of the correct molecule weight composed of two H chains and two L chains was confirmed. Further, the N terminal amino acid sequences for the H and L chains of the purified antihuman IL-5R α human chimeric antibody KM1399 were analyzed with a protein sequencer (470A, Applied Biosystems) by the automatic Edman method. As a result, the expected correct amino acid sequences were obtained.

(4) Reactivity of the Anti-Human IL-5R α Human Chimeric Antibody KM1399 with Human IL-5R α (ELISA method 1)

The reactivities of the anti-human IL-5R α mouse antibody KM1259 and the anti-human IL-5R α human chimeric antibody KM1399 with human IL-5R α were determined by the ELISA method 1 described in subsection (2) of section 3 of Example 2. The results are shown in Fig. 40. As seen from Fig. 40, the anti-human IL-5R α human chimeric antibody KM1399 proved to have a strong reactivity with human IL-5R α which was comparable to the reactivity of the anti-human IL-5R α mouse antibody KM1259.

 σ 4. Transient Expression of Anti-Human IL-5R α Human Chimeric Antibody in COS-7 Cells (ATCC CRL1651)

In order to evaluate the activities of various versions of the anti-human IL-5R α human CDR-grafted antibody to be described later more quickly, the transient expression of an anti-human IL-5R α human chimeric antibody in COS-7 cells was examined as follows using pKANTEX1259 and a modified vector thereof by the lipofectamine method.

(1) Construction of a improved vector of pKANTEX1259

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Since the efficiency of the transient expression of a gene in animal cells depends on the number of copies of the expression vector transfected thereinto, it was assumed that a smaller expression vector would lead to a better expression efficiency. Therefore, a smaller anti-human IL-5R α human chimeric antibody expression vector, pT1259, was constructed as follows by deleting some regions of pKANTEX1259 which were believed not to influence the expression of an antibody.

Briefly, 2 μ g of plasmid pKANTEX1259 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 50 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Hindlll (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Mlul (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated and the 5' sticky ends generated by the digestion with the restriction enzyme were blunted using DNA Blunting Kit (Takara Shuzo). The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 9.60 kb DNA fragment. Then, 0.1 μ g of the recovered DNA fragment was added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to thereby recover plasmid pT1259 shown in Fig. 41.

(2) Transient Expression of Anti-Human IL-5R α Human Chimeric Antibody using pT1259

COS-7 cells at a concentration of 1x10⁵ cells/ml were dispersed into a 6-well plate (2 ml/well) and cultured at 37 °C overnight. To 100 μl of OPTI-MEM (Gibco), 2 μg of pT1259 was added, followed by addition of a solution obtained by adding 10 μ l of lipofectamine reagent (Gibco) to 100 μ l of OPTI-MEM medium (Gibco). The resultant mixture was reacted at room temperature for 40 minutes to thereby form a DNA-liposome complex. COS-7 cells described above were washed with 2 ml of OPTI-MEM medium (Gibco) twice, and the solution containing the DNA-liposome complex was added thereto. Then, the cells were cultured at 37 °C for 7 hours. After the removal of the cultured fluid, 2 ml of DMEM medium (Gibco) containing 10% FCS was added and the cells were cultured at 37°C. At 72 hours from the start of the cultivation, the culture supernatant was recovered, and the activity of an anti-human IL-5R α human chimeric antibody in the culture supernatant was evaluated by the ELISA method 1 described in subsection (2) of section 3 of Example 2. As shown in Fig. 42, concentration-dependent activity was observed in the culture supernatant of COS-7 cells into which pT1259 had been transfected. Thus, the expression of an anti-human IL-5R α human chimeric antibody was confirmed. From these results, it has been shown to be possible to evaluate the activities of humanized antibodies derived from various expression vectors in a transient expression system by preparing a improved small-size vector pKANTEX93, and by then transfecting the vector into COS-7 cells. Further, in order to compare correctly the activities of the various anti-human IL-5R α human CDR-grafted antibodies to be described later, the concentration of antibody formed by the transient expression in culture supernatant was determined by the ELISA method described in subsection (3) of section 4 below.

(3) Determination of the Humanized Antibody Concentration in the Transient Expression-Culture Supernatant by ELISA

To a 96-well microtiter plate, a solution obtained by diluting goat anti-human $IgG(\gamma\text{-chain})$ antibody (Institute of Medicine & Biology) to 400 fold with PBS was dispensed (50 μ I/well) and reacted at 4°C overnight. After the removal of the antibody solution, 100 μ I/well of 1% BSA-PBS was added and reacted at 37°C for 1 hour to thereby block the remaining active groups. After discarding 1% BSA-PBS, 50 μ I/well of the transient expression-culture supernatant or the purified anti-human IL-5R α human chimeric antibody KM1399 was added and reacted at room temperature for 1 hour. After the reaction, the mixture was removed and the plate was washed with 0.05% Tween-PBS. Then, 50 μ I/well of a solution obtained by diluting peroxidase-labeled mouse anti-human κ L chain antibody (Zymed) 500 folds with 1% BSA-PBS was added to the plate and reacted at room temperature for 1 hour. After washing with 0.05% Tween-PBS, 50 μ I/well of ABTS substrate solution [as obtained by dissolving 550 mg of 2,2'azinobis(3-ethylbenzothiazoline-6-sulfonic acid)diammonium in 1 L of 0.1 M citrate buffer (pH 4.2) and adding 1 μ I/ml of hydrogen peroxide immediately before use] was added to allow color development. Then, the absorbance at OD of 415 nm was measured.

5. Preparation of an Anti-Human IL-5Rα Human CDR-Grafted Antibody

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An anti-human IL-5R α human CDR-grafted antibody was prepared as described below; the antibody had a comparable activity to the mouse anti-human IL-5R α monoclonal antibody KM1259 and the anti-human IL-5R α human chimeric antibody KM1399, both of which had an activity to inhibit the biological activity of human IL-5.

(1) Construction of a cDNA coding for the VH of an Anti-Human IL-5R α Human CDR-Grafted Antibody based on the Consensus Sequence for the VH of Known Human Antibodies

Kabat et al. (Sequences of Proteins of Immunological Interest, US Dept. Health and Human Services, 1991) classified various known human antibody VH into subgroups 1-III (HSG I-III) based on the homology of FR sequence, and identified the consensus sequence for each subgroup. The present inventors therefore decided to design an amino acid sequence for an anti-human IL-5R α human CDR-grafted antibody VH based on those consensus sequences. First, in order to select a consensus sequence to be used as the base, the homology between the FR sequence for the VH of the mouse anti-human IL-5R α monoclonal antibody KM1259 and the FR sequence of the consensus sequence of human antibody VH of each subgroup was examined (Table 1).

Table 1

Homology (%) between the FR Sequence for Mouse KM1259VH and the FR Sequence of the Consensus Sequence of Human Antibody VH of Each Subgroup													
HSGI	HSGII	HSGIII											
72.1	72.1 50.6 55.2												

As a result, it was confirmed that mouse KM1259VH has the highest homology to subgroup I in FR sequence. Thus, the amino acid sequence for an anti-human IL-5R α human CDR-grafted antibody VH was designed based on the consensus sequence of subgroup I, and a cDNA coding for the above amino acid sequence was constructed as described below using PCR.

Briefly, 6 synthetic DNAs having the base sequences shown in SEQ ID NOS: 50-55, respectively, were synthesized with an automatic DNA synthesizer (380A; Applied Biosystems). Each of the synthesized DNAs was added to 50 μ l of a buffer containing 10 mM Tris-HCl (pH 8.3), 50 mM potassium chloride, 1.5 mM magnesium chloride, 0.001% gelatin, 200 μ M dNTP, 0.5 μ M M13primer RV (Takara Shuzo), 0.5 μ M M13primer M4 (Takara Shuzo) and 2 units of TaKaRa Taq DNA polymerase (Takara Shuzo) to give a final concentration of 0.1 μ M. Then, the resultant mixture was covered with 50 μ l of mineral oil and set in a DNA thermal cycler (PJ480; Perkin Elmer). Then, PCR was performed through 30 cycles, each cycle consisting of 94°C for 2 minutes, 55°C for 2 minutes and 72°C for 2 minutes. The reaction mixture was ethanol-precipitated and the precipitate was dissolved in 20 μ l of TE buffer. Thereafter, the mixture was subjected to agarose gel electrophoresis to thereby recover about 0.2 μ g of an approx. 0.48 kb amplified fragment.

Subsequently, 3 μ g of plasmid pBluescriptSK(-) (Stratagene) was added to 10 μ l of a buffer containing 33 mM Tris-HCl (pH 7.9), 10 mM magnesium acetate, 66 mM potassium acetate, 0.5 mM DTT and 100 μ g/ml BSA, to which 10 units of the restriction enzyme Smal (Takara Shuzo) was added and reacted at 30 °C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 20 μ l of a buffer containing 50 mM Tris-HCl (pH 9.0) and 1

mM magnesium chloride, to which 1 unit of alkaline phosphatase (E. coli C75, Takara Shuzo) was added and reacted at 37 $^{\circ}$ C for 1 hour to thereby dephosphorylate 5' ends. Then, the reaction mixture was subjected to phenol-chloroform extraction, followed by ethanol precipitation. The precipitate was dissolved in 20 μ l of TE buffer.

Subsequently, 0.1 μ g of the amplified fragment obtained by PCR and 0.1 μ g of the Smal fragment from pBlue-scriptSK(-) were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed. From 10 transformant clones, plasmid DNA was prepared individually and the base sequence thereof was determined. As a result, plasmid phKM1259HV0 shown in Fig. 43 comprising a cDNA coding for the amino acid sequence for an anti-human IL-5R α human CDR-grafted antibody VH of interest was obtained. The base sequence and the amino acid sequence for the anti-human IL-5R α human CDR-grafted antibody VH contained in phKM1259HV0 (hereinafter referred to as " HV.0") are shown in SEQ ID NO: 56.

(2) Construction of a cDNA coding for the VL of an Anti-Human IL-5R α Human CDR-Grafted Antibody based on the Consensus Sequence for the VL of Known Human Antibodies

Kabat et al. classified various known human antibody VL into subgroups 1-IV (HSG I-IV) based on the homology of FR sequence, and identified the consensus sequence for each subgroup. The present inventors therefore decided to design an amino acid sequence for an anti-human IL-5R α human CDR-grafted antibody VL based on those consensus sequences. First, in order to select a consensus sequence to be used as the base, the homology between the FR sequence for the VH of the mouse anti-human IL-5R α monoclonal antibody KM1259 and the FR sequence of the consensus sequence of human antibody VL of each subgroup was examined (Table 2).

Table 2

Homology (%) between the FR Sequence for Mouse KM1259VL and the FR Sequence of the Consensus Sequence of Human Antibody VL of Each Subgroup										
HSGI	HSGII	HSGIII	HSGIV							
73.8	57.5	60.0	65.0							

As a result, it was confirmed that mouse KM1259VL has the highest homology to subgroup I in FR sequence. Thus, the amino acid sequence for an anti-human IL-5R α human CDR-grafted antibody VL was designed based on the consensus sequence of subgroup I, and a cDNA coding for the above amino acid sequence was constructed as described below using PCR.

Briefly, 6 synthetic DNAs having the base sequences shown in SEQ ID NOS: 57-62, respectively, were synthesized with an automatic DNA synthesizer (380A; Applied Biosystems). Each of the synthesized DNAs was added to 50 μ l of a buffer containing 10 mM Tris-HCl (pH 8.3), 50 mM potassium chloride, 1.5 mM magnesium chloride, 0.001% gelatin, 200 μ M dNTP, 0.5 μ M M13primer RV (Takara Shuzo), 0.5 μ M M13primer M4 (Takara Shuzo) and 2 units of TaKaRa Taq DNA polymerase (Takara Shuzo) to give a final concentration of 0.1 μ M. Then, the resultant mixture was covered with 50 μ l of mineral oil and set in a DNA thermal cycler (PJ480; Perkin Elmer). Then, PCR was performed through 30 cycles, each cycle consisting of 94 °C for 2 minutes, 55°C for 2 minutes and 72 °C for 2 minutes. The reaction mixture was ethanol-precipitated and the precipitate was dissolved in 20 μ l of TE buffer. Thereafter, the solution was subjected to agarose gel electrophoresis to thereby recover about 0.2 μ g of an approx. 0.43 kb amplified fragment.

Subsequently, 0.1 μ g of the amplified fragment obtained above by PCR and 0.1 μ g of the Smal Fragment from pBluescriptSK(-) obtained in subsection (1) of section 5 of Example 2 were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed. From 10 transformant clones, plasmid DNA was prepared individually and the base sequence thereof was determined. As a result, plasmid phKM1259LV0 shown in Fig. 44 comprising a cDNA coding for the amino acid sequence for the anti-human IL-5R α human CDR-grafted antibody VL of interest was obtained. The base sequence and the amino acid sequence for the anti-human IL-5R α human CDR-grafted antibody VL contained in phKM1259LV0 (hereinafter referred to as "LV.0") are shown in SEQ ID NO: 63.

(3) Construction of Expression Vector for Anti-Human IL-5R α Human CDR-Grafted Antibody pKANTEX1259HV0LV0, based on the Consensus Sequence of V Regions of Known Human Antibodies

An anti-human IL-5R α human CDR-grafted antibody expression vector, pKANTEX1259HV0LV0, was constructed

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as described below using the humanized antibody expression vector pKANTEX93 obtained in section 1 of Example 2, the plasmid phKM1259HV0 obtained in subsection (1) of section 5 of Example 2 and the plasmid phKM1259LV0 obtained in subsection (2) of section 5 of Example 2.

Briefly, 5 μ g of plasmid pKMh1259HV0 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT, 100 μ g/ml BSA and 0.01% Triton X-100, to which 10 units or the restriction enzyme Notl (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.5 μ g of an approx. 0.44 kb Apal-Notl fragment.

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Subsequently, 0.1 μ g of the Apal-Notl fragment from the humanized antibody expression vector pKANTEX93 obtained in subsection (1) of section 3 of Example 2 and 0.1 μ g of the Apal-Notl fragment from plasmid phKM1259HV0 obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to thereby obtain plasmid pKANTEX1259HV0 shown in Fig. 45.

Subsequently, 3 μ g of the thus obtained plasmid pKANTEX1259HV0 was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT and 100 μ g/ml BSA, to which 10 units each of the restriction enzyme EcoRl (Takara Shuzo) and the restriction enzyme Spll (Takara Shuzo) were added and reacted at 37°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 13.20 kb EcoRl-Spll fragment.

Subsequently, 5 μ g of plasmid phKM1259LV0 was added to 10 μ l of a buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM sodium chloride, 10 mM magnesium chloride, 1 mM DTT and 100 μ g/ml BSA, to which 10 units each of the restriction enzyme EcoRl (Takara Shuzo) and the restriction enzyme Spll (Takara Shuzo) were added and reacted at 37 °C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.5 μ g of an approx. 0.39 kb EcoRl-Spll fragment.

Then, 0.1 μ g of the EcoRI-SpII fragment from plasmid pKANTEX1259HV0 obtained above and 0.1 μ g of the EcoRI-SpII fragment from plasmid phKM1259LV0 obtained above were added to sterilized water to give a total volume of 20 μ I and ligated using Ready-To-Go T4'! DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed to thereby obtain plasmid pKANTEX1259HV0LV0 shown in Fig. 46.

(4) Expression of an Anti-Human IL-5R α Human CDR-Grafted Antibody based on the Consensus Sequence of Known Human Antibody V Regions in Rat Myeloma YB2/0 Cells (ATCC CRL1581) using pKANTEX1259HV0LV0

The expression of an anti-human IL-5R α human CDR-grafted antibody based on the consensus sequence of known human antibody V regions in rat myeloma YB2/0 cells (ATCC CRL1581) was performed using pKANTEX1259HV0LV0 according to the method described in subsection (2) of section 3 of Example 2.

As a result, KM8397 was obtained as a transformant producing an anti-human IL-5R α human CDR-grafted anti-body based on the consensus sequence of known human antibody V regions. The anti-human IL-5R α human CDR-grafted antibody produced by the strain was designated as KM8397. The productivity of the anti-human IL-5R α human CDR-grafted antibody KM8397 in the transformant KM8397 was about 4 μ g/10⁶ cells/24 hr.

(5) Purification of the Anti-Human IL-5R α Human CDR-Grafted Antibody KM8397 from Culture Supernatant

The anti-human IL-5R α human CDR-grafted antibody-producing clone KM8397 obtained in subsection (4) of section 5 of Example 2 was cultured according to the method described in subsection (3) of section 3 of Example 2 and purified to thereby obtain about 2 mg of KM8397. About 4 μ g of the purified anti-human IL-5R α human CDR-grafted antibody KM8397 was electrophoresed according to the method described in subsection (3) of section 3 of Example 2 in order to examine its molecular weight. The results are shown in Fig. 47. As shown in Fig. 47, the molecular weight of the antibody H chain is about 50 KDa and that of the antibody L chain about 25 KDa under reducing conditions. Thus, the expression of the H and L chains with the correct molecular weights was confirmed. On the other hand, under non-reducing conditions, the molecular weight of the anti-human IL-5R α human CDR-grafted antibody KM8397 is about 140 KDa. Thus, the expression of a human CDR-grafted antibody of the correct size composed of two H chains and two L chains was confirmed. Further, the N terminal amino acid sequences for the H and L chains of the purified anti-human IL-5R α human CDR-grafted antibody KM8397 were analyzed with a protein sequencer (470A, Applied Biosystems) by the automatic Edman method. As a result, the correct amino acid sequences as expected were obtained.

(6) Reactivity of the Anti-Human IL-5R α Human CDR-Grafted Antibody KM8397 with Human IL-5R α (ELISA method 2)

The reactivities of the anti-human IL-5R α human chimeric antibody KM1399 and the anti-human IL-5R α human CDR-grafted antibody KM8397 with human IL-5R α were determined by the ELISA method 2 described in subsection (2) of section 3 of Example 2. The results are shown in Fig. 48. As shown in Fig. 48, the reactivity of the anti-human IL-5R α human CDR-grafted antibody KM8397 with human IL-5R α was shown to be about one half the reactivity of the anti-human IL-5R α human chimeric antibody KM1399.

6. Increase in Activity by Modification of the Amino Acid Sequence for the V Region of the Anti-Human IL-5R α Human CDR-Grafted Antibody KM8397

The reactivity of the anti-human IL-5R α human CDR-grafted antibody KM8397 with human IL-5R α decreased to about one half the reactivity of the anti-human IL-5R α human chimeric antibody KM1399.

Therefore, the activity of KM8397 was increased by modifying the amino acid sequence for the V region thereof by the methods described below.

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(1) Modification of the Amino Acid Sequence for VH of the Anti-Human IL-5R α Human CDR-Grafted Antibody KM8397

By mutating the amino acids of VH of the anti-human IL-5R α human CDR-grafted antibody KM8397 shown in SEQ ID NO: 56, various modified versions of VH of the anti-human IL-5R α human CDR-grafted antibody were prepared. The amino acids to be mutated were selected at random with reference to a computerized three-dimensional structural model for the V region of the anti-human IL-5R α mouse antibody KM1259. As the method for transfecting a mutation, a plasmid comprising a cDNA coding for a modified version of VH of interest of the anti-human IL-5R α human CDR-grafted antibody was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using primers for mutation.

Actually, a plasmid, phKM1259HV1, comprising a cDNA coding for the modified version 1 of VH (hereinafter referred to as "HV.1") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 65 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequence shown in SEQ ID NO: 64 as a primer for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 50, 51, 52, 53, 64 and 55, respectively. In the amino acid sequence of HV.1, tyrosine in position 95 and alanine in position 97 located in the FR of SEQ ID NO: 56 have been replaced with leucine and glycine, respectively, which are the amino acids found in the V region of the mouse antibody KM1259 H chain and this is in order to retain the reactivity with human IL-5R α recognized in the mouse antibody and the human chimeric antibody.

Further, a plasmid, phKM1259HV2, comprising a cDNA coding for the modified version 2 of VH (hereinafter referred to as "HV.2") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 68 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NOS: 64, 66 and 67 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 50, 51, 66, 67, 64 and 55, respectively. In the amino acid sequence of HV.2, glutamic acid in position 46, threonine in position 74, tyrosine in position 95 and alanine in position 97 located in the FR of SEQ ID NO: 56 have been replaced with alanine, arginine, leucine and glycine, respectively, which are the amino acids found in the V region of the mouse antibody KM1259 H chain and this is in order to retain the reactivity with human IL-5R α recognized in the mouse antibody and the human chimeric antibody.

Further, a plasmid, phKM1259HV3, comprising a cDNA coding for the modified version 3 of VH (hereinafter referred to as "HV.3") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 72 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NOS: 69, 70 and 71 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 50, 51, 69, 70, 71 and 55, respectively. In the amino acid sequence of HV.3, alanine in position 40, glutamic acid in position 46, arginine in position 67, alanine in position 72, threonine in position 74, alanine in position 79, tyrosine in position 95 and alanine in position 97 located in the FR of SEQ ID NO: 56 have been replaced with arginine, alanine, lysine, serine, arginine, valine, leucine and glycine, respectively, which are the amino acids found in the V region of the mouse antibody KM1259 H chain and this is in order to retain the reactivity with human IL-5R α recognized in the mouse antibody and the human chimeric antibody.

As version advances from HV.0 to HV.4 one by one, the number of the monoclonal antibody-derived amino acids involved in the modification increases with increasing version number from HV.0 to HV. 3.

(2) Modification of the Amino Acid sequence for VL of the Anti-Human IL-5R lpha Human CDR-Grafted Antibody KM8397

By mutating the amino acids of VL of the anti-human IL-5R α human CDR-grafted antibody KM8397 shown in SEQ

ID NO: 63, various modified versions of VL of the anti-human IL-5R α human CDR-grafted antibody were prepared. The amino acids to be mutated were selected at random with reference to a computerized 3D structural model for the V region of the anti-human IL-5R α antibody KM1259. As the method for transfecting a mutation, a plasmid comprising a cDNA coding for a modified version of VL of interest of the anti-human IL-5R α human CDR-grafted antibody was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using primers for mutation.

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Actually, a plasmid, phKM1259LV1, comprising a cDNA coding for the modified version 1 of VL (hereinafter referred to as "LV.1") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 76 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NO: 73, 74 and 75 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 57, 58, 73, 74, 61 and 75, respectively. In the amino acid sequence of LV.1, glutamine in position 37, lysine in position 45 and phenylalanine in position 98 located in the FR of SEQ ID NO: 63 have been replaced with arginine, glutamic acid and valine, respectively, which are the amino acids found in the V region of the monoclonal antibody KM1259 L chain and this is in order to retain the reactivity with human IL-5R α recognized in the monoclonal antibody and the human chimeric antibody.

Further, a plasmid, phKM1259LV2, comprising a cDNA coding for the modified version 2 of VL (hereinafter referred to as "LV.2") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 79 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NOS: 74, 75, 77 and 78 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 57, 58, 77, 74, 78 and 75, respectively.

In the amino acid sequence for LV.2, threonine in position 22, glutamine in position 37, lysine in position 45, serine in position 77 and phenylalanine in position 98 located in the FR of SEQ ID NO: 63 have been replaced with glycine, arginine, glutamic acid, aspartic acid and valine, respectively, which are the amino acids found in the V region of the monoclonal antibody KM1259 L chain and this is in order to retain the reactivity with human IL-5R α recognized in the monoclonal antibody and the human chimeric antibody.

Further, a plasmid, phKM1259LV3, comprising a cDNA coding for the modified version 3 of VL (hereinafter referred to as "LV.3") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 84 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NOS: 74, 80, 81, 82 and 83 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 57, 80, 81, 74, 82 and 83, respectively. In the amino acid sequence of LV.3, serine in position 7, proline in position 8, threonine in position 22, glutamine in position 37, glutamine in position 38, lysine in position 45, serine in position 77, tyrosine in position 87 and phenylalanine in position 98 located in the FR of SEQ ID NO: 63 have been replaced with alanine, threonine, glycine, arginine, lysine, glutamic acid, aspartic acid, phenylalanine and valine, respectively, which are the amino acids found in the V region of the monoclonal antibody KM1259 L chain and this is in order to retain the reactivity with human IL-5R α recognized in the monoclonal antibody and the human chimeric antibody.

Further, a plasmid, phKM1259LV4, comprising a cDNA coding for the modified version 4 of VL (hereinafter referred to as "LV.4") of the anti-human IL-5R α human CDR-grafted antibody shown in SEQ ID NO: 88 was obtained by performing the procedures described in subsection (1) of section 5 of Example 2 using the sequences shown in SEQ ID NOS: 80, 83, 85, 86 and 87 as primers for mutation and using synthetic DNAs having base sequences of SEQ ID NOS: 57, 80, 85, 86, 87 and 83, respectively. In the amino acid sequence of LV.4, serine in position 7, proline in position 8, threonine in position 22, glutamine in position 37, glutamine in position 38, proline in position 44, lysine in position 45, phenylalanine in position 71, serine in position 77, tyrosine in position 87 and phenylalanine in position 98 located in the FR of SEQ ID NO: 63 have been replaced with alanine, threonine, glycine, arginine, lysine, valine, glutamic acid, tyrosine, aspartic acid, phenylalanine and valine, respectively, which are the amino acids found in the V region of the monoclonal antibody KM1259 L chain and this is in order to retain the reactivity with human IL-5R α recognized in the monoclonal antibody and the human chimeric antibody.

As a result, as version advances from LV.0 to HV.4 one by one, the number of the monoclonal antibody-derived amino acids involved in the modification increases with increasing version number from LV.0 to LV.4.

(3) Preparation of Anti-Human IL-5R α Human CDR-Grafted Antibodies having Various Modified Versions of V Region

Using the humanized antibody expression vector pKANTEX93 constructed in section 1 of Example 2 and the various plasmids comprising cDNAs coding for various modified versions of the V region of the anti-human IL-5R α human CDR-grafted antibody obtained in subsections (1) and (2) of section 5 of Example 2, vectors for the expression of anti-human IL-5R α human CDR-grafted antibodies having various modified versions of the V region were constructed by the method described in subsection (3) of section 5 of Example 2. Table 3 shows combinations of various modified versions of the V region used in the expression vectors constructed and the designation of these expression vectors.

Table 3

5	YL VI	H HY. 0	HV. 1	HV. 2	HV. 3
	LY. 0	pKANTEX1259HV0LV0	pKANTEX1259HY1LY0	pKANTEX1259HV2LY0	pKANTEX1259HV3LVO
	LY. I	pKANTEX1259HVOLV1	pKANTEX1259HV1LV1	pKANTEX1259HY2LY1	
40	LY. 2	pKANTEX 1 259HVOLV2	pKANTEX1259HV1LV2	pKANTEX1259HY2LY2	
10	LY. 3	pKANTEX1259HVOLV3	pKANTEX1259HV1LV3	pKANTEX1259HV2LV3	pKANTEX1259HY3LY3
	LY. 4	pKANTEX I 259HVOLV4	pKANTEX1259HV1LV4	pKANTEX1259HY2LY4	pKANTEX1259HV3LV4

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Among these expression vectors, a total of 13 vectors pKANTEX1259HV0LV0, pKANTEX1259HV1LV0, pKANTEX1259HV2LV0, pKANTEX1259HV0LV1, pKANTEX1259HV1LV1, pKANTEX1259HV2LV1, pKANTEX1259HV0LV2. pKANTEX1259HV1LV2, pKANTEX1259HV2LV2, pKANTEX1259HV0LV3. pKANTEX1259HV1LV3, pKANTEX1259HV2LV3, and pKANTEX1259HV3LV3 were modified into transient expression vectors by the method described in subsection (1) of section 4 of Example 2. Using these transient expression vectors and in accordance with the method described in subsection (2) of section 4 of Example 2, the transient expression of anti-human IL-5R α human CDR-grafted antibodies having various modified versions of the V region was performed. As a control, the transient expression of the anti-human IL-5R α human chimeric antibody KM1399 was performed simultaneously. The binding activity for human IL-5R α of an antibody in the culture supernatant was determined by the ELISA method 1 described in subsection (2) of section 3 of Example 2, and the antibody concentration in the culture supernatant was determined by the ELISA method described in subsection (3) of section 4 of Example 2. Using two ELISA methods, the activities of anti-human IL-5R α human CDR-grafted antibodies having various modified versions of the V region are shown in Fig. 49 as relative values in which the activity of the human chimeric antibody KM1399 is taken as 100. In Fig. 49, various modified versions of anti-human IL-5R α human CDR-grafted antibodies are represented by a combination of VH and VL. From Fig. 49, a tendency is recognized with VH such that the activity increases as modification proceeds from HV.0 to HV.3. With respect to VL, a tendency is recognized such that the reactivity is high in LV.0 and LV.3 but low in LV.1 and LV.2. Then, a more accurate activity evaluation of anti-human IL-5R α human CDRgrafted antibodies comprising combinations of LV.0 and various modified VH; LV.3 and HV.0; LV.3 and HV.3; and LV.4 which is a further modified version of LV.3, and various modified VH was performed using purified antibodies as follows.

Briefly, using the 10 expression vectors for anti-human IL-5R α human CDR-grafted antibodies described above, pKANTEX1259HV0LV0, pKANTEX1259HV1LV0, pKANTEX1259HV2LV0, pKANTEX1259HV3LV0, pKANTEX1259HV3LV3, pKANTEX1259HV0LV3, pKANTEX1259HV0LV4, pKANTEX1259HV1LV4, pKANTEX1259HV2LV4, and pKANTEX1259HV3LV4 and according to the method described in subsection (2) of section 3 of Example 2, antibodies of interest were expressed in YB2/0 cells to thereby obtain transformant producing various anti-human IL-5R α human CDR-grafted antibodies at a productivity level of 2-4 μ g/10⁶ cells/24 hr. The transformants producing various anti-human IL-5R α human CDR-grafted antibodies were cultured and purified by the methods described in subsection (3) of section 3 of Example 2 to thereby obtain 1-2 mg each of various anti-human IL- $5R \alpha$ human CDR-grafted antibodies. About 4 μg each of the various purified anti-human IL- $5R \alpha$ human CDR-grafted antibodies were electrophoresed by the method described in subsection (3) of section 3 of Example 2 to measure their molecular weights. Under reducing conditions, the molecular weight of the antibody H chain is about 50 kDa and that of the antibody L chain about 25 kDa in each of the anti-human IL-5R α human CDR-grafted antibodies. Thus, the expression of H and L chains with the correct molecular weights was confirmed. Under non-reducing conditions, the molecular weight of the antibody is about 140 kDa in each of the anti-human IL-5R α human CDR-grafted antibodies. Thus, the expression of human CDR-grafted antibodies each composed of two H chains and two L chains of the correct size was confirmed. Further, the N terminal amino acid sequences for the H and L chains of the various purified antihuman IL-5R α human CDR-grafted antibodies were analyzed with a protein sequencer (470A, Applied Biosystems) by the automatic Edman method. As a result, the correct amino acid sequences as expected were obtained in each of those antibodies.

The reactivity with human IL-5R α in the various purified anti-human IL-5R α human CDR-grafted antibodies obtained above was determined by the ELISA method 2 described in subsection (2) of section 3 of Example 2 and the results are shown in Fig. 50. In Fig. 50, various modified versions of anti-human IL-5R α human CDR-grafted antibodies are represented by a combination of VH and VL. As shown in Fig. 50, of the 10 purified anti-human IL-5R α human CDR-grafted antibodies, HV.3LV.0 and HV.3LV.4 proved to have a reactivity with human IL-5R α as strong as the reac-

tivity of the anti-human IL-5R α human chimeric antibody KM1399.

When the amino acid sequences for the anti-human IL-5R α human CDR-grafted antibodies HV.3LV.0 and HV.3LV.4 exhibiting a reactivity with human IL-5R α as strong as the reactivity of the anti-human IL-5R α human chimeric antibody KM1399 are compared, both have the same amino acid sequence which is shown as HV.3 for VH but they have different amino acid sequences for VL, i.e., shown as LV.0 and LV.4. While LV.0 is a sequence obtained by simply grafting the CDR to the FR of a human antibody, LV.4 is a sequence obtained by converting 11 amino acid residues within the FR of a human antibody to those amino acid residues found in the monoclonal antibody in order to increase activity. However, from the results shown in Fig. 50, the modification of amino acid residues makes little contributions to the increase of activity actually. Based on these facts, HV.3LV.0 which has a reactivity with human IL-5R α as strong as the reactivity of the anti-human IL-5R α human chimeric antibody KM1399 and which is expected to be less antigenic against humans since the replacement of amino acids derived from the monoclonal antibody is less, has been selected as an anti-human IL-5R α human CDR-grafted antibody. HV.3LV.0 was designated as KM8399, and the transformant KM8399 producing the anti-human IL-5R α human CDR-grafted antibody KM8399 was deposited with the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on September 3, 1996 under accession number FERM BP-5648.

In the preparation of the anti-human IL-5R α human CDR-grafted antibody KM8399, the following matters have been taken into consideration. As seen in the preparation of other human CDR-grafted antibodies, the activity in the anti-human IL-5R α human CDR-grafted antibody KM8397, which was obtained by simply grafting only the CDR of the anti-human IL-5R α monoclonal antibody KM1259 into the FR of a human antibody, decreased to about 1/2 of the activity of the monoclonal antibody KM1259. Hence, several amino acids within the FR of the V regions of H and L chains were modified into the amino acids found in the monoclonal antibody KM1259, and examined for an increase in activity. With respect to VH, the activity increased as the modification proceeded. On the other hand, with respect to VL, the modification of a small number of amino acids resulted in a decrease in activity; although the activity can be increased by increasing the number of amino acids modified, the activity only rose to the level of unmodified VL. Although the cause of this fact cannot be completely clarified without more detailed analysis (e.g., X-ray crystal analysis), the interaction between the VH and VL of an antibody is probably be involved and the results of such interaction would vary depending on the antibody used. Because of such problems, no efficient method has yet been established for preparing a human CDR-grafted antibody of which is applicable to any antibody and trials and errors as made in the present Example are required. With such trials and errors being accumulated, a more efficient method for preparing human CDR-grafted antibodies could be established. The present Example shows the first case of successful preparation of an anti-human IL-5R α human CDR-grafted antibody and thus provides suggestions for efficient preparation of human CDR-grafted antibodies.

7. Preparation of Anti-Human IL-5R lpha Humanized Antibodies of Human Antibody IgG4 Subclass

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(1) Isolation and Analysis of a cDNA coding for the C Region ($C\gamma4$) of Human Antibody IgG4 Subclass

1.1x10⁷ B cells were separated from 200 ml of peripheral blood from a healthy volunteer using anti-CD19 antibody coated Dynabeads (DYNABEADS M-450 Pan-B(CD19); Nippon Dyner) and DETACHaBEAD (Nippon Dyner) in accordance with the attached instructions. Then, mRNA was obtained from the separated cells using Quick Prep mRNA Purification Kit (Pharmacia Biotech) in accordance with the attached instructions. From all of the mRNA obtained, cDNA was synthesized using Time Saver cDNA Synthesis Kit (Pharmacia Biotech) in accordance with the attached instructions. Then PCR was performed as described in subsection (1) of section 5 of Example 2 using all of the cDNA obtained above and using, as primers, synthetic DNAs shown in SEQ ID NOS: 89 and 90 which are homologous to the 5' and 3' sides of a cDNA coding for human antibody Cγ4 [Nucleic Acid Research, 14, 1789 (1986)]. The 5' side and 3' side primers used in the PCR had been designed to have recognition sequences for the restriction enzymes Apal and BamHI at their 5' terminals so that the cDNA to be obtained could be easily inserted into a humanized antibody expression vector. The reaction mixture after the PCR was purified with QIAquick PCR Purification Kit (Qiagen) and then added to 30 μl of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT. To the resultant mixture, 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated and the precipitate was added to 10 µl of a buffer containing 20 mM Tris-HCl (pH 8.5), 100 mM potassium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme BamHI (Takara Shuzo) was added and reacted at 30°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 0.5 µg of an approx. 1.0 kb Apal-BamHI fragment.

Subsequently, 3 μ g of plasmid pBluescriptSK(-) (Stratagene) was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 100 mM potassium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme BamHl (Takara Shuzo) was added and reacted at

 30° C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 2 μ g of an approx. 3.0 kb Apal-BamHI fragment.

Then, 0.1 μ g of the PCR-amplified Apal-BamHI fragment obtained above and 0.1 μ g of the Apal-BamHI fragment from pBluescriptSK(-) obtained above were added to sterilized water to give a total volume of 20 μ l and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using the thus obtained recombinant plasmid DNA solution, E. coli HB101 was transformed. From 10 transformant clones, each plasmid DNA was prepared and the base sequence thereof was determined.

As a result, plasmid pBShC γ 4 shown in Fig. 51 comprising a cDNA of interest coding for human antibody C γ 4 was obtained.

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(2) Construction of an Expression Vector for Anti-Human IL-5R α Humanized Antibodies of Human Antibody IgG4 Subclass

An expression vector for anti-human IL-5R α humanized antibodies of human antibody IgG4 subclass was constructed as described below using plasmid pBShC γ 4 comprising a cDNA coding for human antibody C γ 4 obtained in subsection (1) of section 7 of Example 2, expression vector pKANTEX1259 for the anti-human IL-5R α human chimeric antibody KM1399 obtained in subsection (1) of section 3 of Example 2 and expression vector pKANTEX1259HV3LV0 for the anti-human IL-5R α human CDR-grafted antibody KM8399 obtained in subsection (3) of section 6 of Example 2.

Briefly, 4 μ g of plasmid pBShC γ 4 comprising a cDNA coding for human antibody C γ 4 was added to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. The reaction mixture was ethanol-precipitated, and the precipitate was added to 10 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 100 mM potassium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme BamHl (Takara Shuzo) was added and reacted at 30°C for 1 hour. The reaction mixture was subjected to agarose gel electrophoresis to thereby recover about 1 μ g of an approx. 1.0 kb Apal-BamHl fragment.

Subsequently, 3 μ g each of expression vector pKANTEX1259 for the anti-human IL-5R α human type chimeric antibody KM1399 and expression vector pKANTEX1259HV3LV0 for the anti-human IL-5R α human CDR-grafted antibody KM8399 were added individually to 10 μ l of a buffer containing 10 mM Tris-HCl (pH 7.5), 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme Apal (Takara Shuzo) was added and reacted at 37°C for 1 hour. Both reaction mixtures were ethanol-precipitated, and the precipitates were individually added to 10 μ l of a buffer containing 20 mM Tris-HCl (pH 8.5), 100 mM potassium chloride, 10 mM magnesium chloride and 1 mM DTT, to which 10 units of the restriction enzyme BamHl (Takara Shuzo) was added and reacted at 30°C for 1 hour. Both reaction mixtures were subjected to agarose gel electrophoresis to thereby recover about 2 μ g of an approx. 12.59 kb Apal-BamHl fragment from each reaction mixture.

A combination of 0.1 μg of the Apal-BamHI fragment from plasmid pBShC $\gamma 4$ and 0.1 μg of the Apal-BamHI fragment from plasmid pKANTEX1259 and another combination of 0.1 μg of the Apal-BamHI fragment from plasmid pBShC $\gamma 4$ and 0.1 μg of the Apal-BamHI fragment from plasmid pKANTEX1259HV3LV0 were added individually to sterilized water to give a total volume of 20 μl and ligated using Ready-To-Go T4 DNA Ligase (Pharmacia Biotech). Using each of the thus obtained recombinant plasmid DNA solutions, E. coli HB101 was transformed to thereby obtain expression vector pKANTEX1259 $\gamma 4$ for an anti-human IL-5R α human chimeric antibody of IgG4 subclass and expression vector pKANTEX1259HV3LV0 $\gamma 4$ for an anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass shown in Fig. 52

(3) Expression of Anti-Human IL-5R α Humanized Antibodies in Rat Myeloma YB2/0 Cells (ATCC CRL1581)

The expression of anti-human IL-5R α humanized antibodies in YB2/0 Cells was performed by the method described in subsection (2) of section 3 of Example 2 using the expression vector pKANTEX1259 γ 4 for an anti-human IL-5R α human chimeric antibody of IgG4 subclass and the expression vector pKANTEX1259HV3LV0 γ 4 for an anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass obtained in subsection (2) of section 7 which were obtained in Example 2.

As a result, as a transformant producing an anti-human IL-5R α human chimeric antibody of IgG4 subclass, KM7399 (FERM BP-5649) was obtained and the anti-human IL-5R α human chimeric antibody of IgG4 subclass produced by this strain was designated as KM7399. The transformant KM7399 producing the anti-human IL-5R α human chimeric antibody KM7399 was deposited with the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on September 3, 1996 under accession number FERM BP-5649. The productivity of the anti-human IL-5R α human chimeric antibody KM7399 in the transformant KM7399 was about 3 μ g/10⁶ cells/24 hr.

Also, as a transformant producing an anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass, KM9399 (FERM BP-5647) was obtained and the anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass produced

by this strain was designated as KM9399. The productivity of the anti-human IL-5R α human CDR-grafted antibody KM9399 in the transformant KM9399 was about 7 μ g/10⁶ cells/24 hr. The transformant KM9399 producing the anti-human IL-5R α human CDR-grafted antibody KM9399 was deposited with the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology on September 3, 1996 under accession number FERM BP-5647.

(4) Purification of the Anti-Human IL-5R α Humanized Antibodies of Human Antibody IgG4 Subclass from Culture Supernatants

The transformant KM7399 producing the anti-human IL-5R α human chimeric antibody of IgG4 subclass and the transformant KM9399 producing the anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass which were obtained in subsection (3) of section 7 of Example 2 were cultured and purified according to the methods described in subsection (3) of section 3 of Example 2, to thereby obtain about 1 mg of KM7399 and about 5 mg of KM9399. About 4 μ g each of the purified anti-human IL-5R α humanized antibodies of IgG4 subclass KM7399 and KM9399 were electrophoresed according to the method described in subsection (3) of section 3 of Example 2 to examine their molecular weights. The results are shown in Fig. 53. As shown in Fig. 53, the molecular weight of the H chain of each antibody is about 50 kDa and that of the L chain of each antibody about 25 kDa under reducing conditions. Thus, the expression of H chains and L chains of the correct molecular weight was confirmed. Under non-reducing conditions, the molecular weight of each anti-human IL-5R α humanized antibody is about 140 kDa. Thus, the expression of a human CDR-grafted antibody of the correct size composed of two H chains and two L chains was confirmed. Further, the N terminal amino acid sequences for the H and L chains of the purified anti-human IL-5R α humanized antibodies of IgG4 subclass KM7399 and KM9399 were analyzed with a protein sequencer (470A, Applied Biosystems) by the automatic Edman method. As a result, the correct amino acid sequences as expected were obtained.

(5) Reactivities of the Anti-Human IL-5R α Humanized Antibodies of Human Antibody IgG4 Subclass with Human IL-5R α (ELISA method 2)

The reactivities of the anti-human IL-5R α human chimeric antibody of human antibody IgG1 subclass KM1399, the anti-human IL-5R α human CDR-grafted antibody of human antibody IgG1 subclass KM8399, the anti-human IL-5R α human chimeric antibody of IgG4 subclass KM7399 and the anti-human IL-5R α human CDR-grafted antibody of IgG4 subclass KM9399 with human IL-5R α were determined by the ELISA method 2 described in subsection (2) of section 3 of Example 2. The results are shown in Fig. 54. As shown in Fig. 54, the anti-human IL-5R α humanized antibodies of human antibody IgG4 subclass proved to have a reactivity with human IL-5R α as strong as the reactivity of the anti-human IL-5R α humanized antibodies of IgG1 subclass.

EXAMPLE 3

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1. Confirmation of the Specificity of Anti-hIL-5R α Antibodies

The specificity of anti-hIL-5R α monoclonal antibodies and anti-hIL-5R α humanized antibodies was confirmed by the following procedures using immunocyte staining.

Briefly, $5x10^5$ cells obtained by transfecting a human IL-5R gene into CTLL-2 cells (ATCC TIB 214) [hereinafter referred to as "CTLL-2(h5R)"] [J. Exp. Med., $\underline{177}$, 1523 (1993)] or $5x10^5$ CTLL-2 cells as a control were suspended in an immunocyte staining buffer (PBS containing 1% BSA, 0.02% EDTA and 0.05% sodium azide) and dispensed into a round bottom 96-well plate (100 μ l/well). After centrifuging at 350xg for 1 minute at 4 °C, the supernatant was discarded. Then, $50~\mu$ l of the immunocyte staining buffer containing 10 μ g/ml of an hIL-5R α antibody was added to each well and reacted at 4 °C for 30 minutes. After the reaction, the immunocyte staining buffer was added (200 μ l/well) and centrifuged at 350xg for 1 minute at 4°C and then the supernatant was removed to wash the cells. The washing operation was further repeated twice. Thereafter, $50~\mu$ l of the immunocyte staining buffer containing FITC-labeled anti-mouse immunoglobulin antibody or FITC-labeled anti-human immunoglobulin antibody (both manufactured by Wako Pure Chemical Industries, Ltd.) diluted 30 folds with a staining buffer was added to each well and reacted at 4°C for 30 minutes. After the reaction, a similar washing operation was repeated three times. Then, the cells were analyzed with a flow cytometer (Coulter).

The results are shown in Fig. 55. Monoclonal antibodies KM1257, KM1259 and KM1486 and humanized antibodies KM1399, KM7399, KM8399 and KM9399 did not react with CTLL-2 cells, but specifically reacted with CTLL-2(h5R). Thus, it has become clear that humanized antibodies KM1399, KM7399, KM8399 and KM9399 specifically recognize hIL-5R α .

2. Action of Anti-IL-5R α Antibodies to Inhibit the Biological Activity of IL-5

Since CTLL-2(h5R) cells exhibit a proliferation response depending on human IL-5 [J. Exp. Med., $\underline{177}$, 1523 (1993)], the effect of the anti-IL-5R α antibodies upon human IL-5 dependent cell proliferation in CTLL-2(h5R) cells was examined. Cell proliferation was evaluated by a color development method using Cell Counting Kit (Dojin Chemical Laboratory).

Briefly, $1x10^4$ CTLL-2(h5R) cells were suspended in 50 μ l of a normal medium and dispensed into a 96-well cell culture plate. These cells were mixed with 25 μ l/well of various anti-IL-5R α antibodies diluted with a normal medium at 40 μ g/ml and with 25 μ l/well of a normal medium containing human IL-5 at 0.4 ng/ml as prepared by the method described in section 3 of Example 1 and cultured in a CO₂ incubator at 37 °C for 44 hours. Then, 10 μ l/well of Cell Counting Kit solution was added to the plate and cells were cultured under 5% CO₂ incubator at 37 °C for another 4 hours. After completion of the cultivation, the absorbance at 450 nm was measured with Microwell Plate Reader Emax (Molecular Device). The CTLL-2(h5R) cell proliferation inhibiting activity of each antibody was calculated by the following formula.

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Percent proliferation inhibition (%) =
$$100 - \frac{A - C}{B - C} \times 100$$

wherein

A: OD value in the presence of an antibody

B: OD value in the absence of an antibody

C: OD value in the absence of human IL-5.

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The results are shown in Fig. 56. Monoclonal antibodies KM1259 and KM1486 and humanized antibodies KM1399, KM8399 and KM9399 inhibited the human IL-5 dependent proliferation of CTLL-2(h5R) cells. However, such activity was not recognized in monoclonal antibody KM1257.

3. Immunocyte Staining of Human Eosinophils

A polymorphonuclear leukocyte fraction was prepared from normal human blood and cultured for 3 days in the presence of human IL-5 to concentrate eosinophils. Then, the reactivity of anti-hIL-5R α monoclonal antibodies was examined with a flow cytometer.

Briefly, polymorphprep (Nicomed) was dispensed into eight 15-ml polypropylene centrifuge tubes (4 ml/tube) and each plated with 6 ml of heparinized normal human blood. Then, the tubes were centrifuged at 500xg for 30 minutes at room temperature to separate and recover polymorphonuclear leukocytes. The polymorphonuclear leukocytes were suspended in a normal medium to give a concentration of 1.25×10^7 cells/10 ml and dispensed into 4 cell culture dishes in 10 ml portions. Then, human IL-5 was added to the cell suspension at a final concentration of 2 ng/ml and the cells were cultured in a CO_2 incubator at 37°C for 3 days. After completion of the cultivation, the cells were centrifuged (1,200 rpm, 5 min.) and suspended in the immunocyte staining buffer to give a concentration of 5×10^6 cells/ml.

Then, 5x10⁵ cells were dispensed into a round bottom 96-well plate.

After the plate was centrifuged at 350xg for 1 minute at 4°C, the supernatant was removed. Then, 50 μ l of 10% normal mouse serum-containing immunocyte staining buffer was added and reacted at 4°C for 30 minutes. To the buffer, monoclonal antibody KM1259 labeled with biotin by conventional methods ["KOSO-KOTAI-HO" (Enzyme Antibody Method), Gakusai Kikaku Co., 1985] or, as a control, biotin-labeled anti-human granulocyte colony-stimulating factor monoclonal antibody KM341 [Agr. Biol. Chem., 53, 1095 (1989)] had been added at a concentration of 10 μ g/ml. After the reaction, 200 μ l of the immunocyte staining buffer was added to each well and centrifuged at 350xg for 1 minutes at 4°C and then the supernatant was removed and the cells were washed. The washing operation was further repeated twice. Thereafter, phycoerythrin-labeled streptavidin (Becton Dickinson) diluted 10 folds with the immunocyte staining buffer was added (50 μ l/well) and reacted at 4°C for 30 minutes. After the reaction, a similar washing operation was repeated 3 times. Then, analysis was performed with a flow cytometer (Coulter) by forward scattering and 90° scattering for those cells which were recognized as polymorphonuclear leukocytes. Also, the same cells were stained by the May-Grunwald-Giemsa staining method ["SENSHOKUHOU NO SUBETE" (Review of Staining Methods), Ishiyaku Shuppan Co., 1988] and observed for polymorphonuclear leukocytes. As a result, it was confirmed that 75% of the cells were eosinophils.

Fig. 57 shows the histogram obtained. Anti-human IL-5R α monoclonal antibody KM1259 exhibited a definite reactivity. Since 75% of the cells analyzed proved to be eosinophils, it was confirmed that anti-human IL-5R α monoclonal antibody KM1259 has a reactivity with human eosinophils.

4. Survival Inhibition of Human Eosinophils with Anti-IL-5R α Antibodies

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Polymorphonuclear leukocyte fractions were prepared from normal human blood, and the action of anti-IL-5R α antibodies upon the survival of eosinophils in the presence of human IL-5 was examined.

Briefly, polymorphprep (Nicomed) was dispensed in 4 ml portions into fifteen 15-ml polypropylene centrifuge tubes and each plated with 8 ml of heparinized normal human blood. Then, the tubes were centrifuged at 500xg for 30 minutes at room temperature to separate and recover polymorphonuclear leukocytes.

Percoll stock solution was prepared by adding 1 volume of sterilized 1.5 M NaCl solution to 9 volumes of Percoll solution (Pharmacia). Then, 80% Percoll solution was prepared by adding 2 volumes of physiological saline to 8 volumes of Percoll stock solution, and 60% Percoll solution was prepared by adding 4 volumes of physiological saline to 6 volumes of Percoll stock solution. For the purpose of removing concomitant monocytes, 5 ml of 60% Percoll solution was dispensed into each of two 15 ml polypropylene centrifuge tubes, plated with the previously obtained polymorphonuclear leukocytes suspended in RPMI1640 medium and centrifuged at 500xg for 30 minutes at room temperature to separate and recover the precipitated polymorphonuclear leukocytes. Further, for the purpose of removing concomitant erythrocytes, 5 ml of 80% Percoll solution was dispensed into each of two 15-ml polypropylene centrifuge tubes, plated with the previously obtained polymorphonuclear leukocytes suspended in RPMI1640 medium and centrifuged at 500xg for 30 minutes at room temperature to separate and recover the polymorphonuclear leukocytes suspended in the Percoll layer.

Subsequently, cells were dispensed into a 48-well cell culture plate at $2x10^6$ cells/well and human IL-5 was added at a final concentration of 0.1 ng/ml. Further, each of various anti-IL-5R α antibodies was added at a final concentration of 1 μ g/ml. For each antibody, 2 wells were cultured and the solution in each well was adjusted to have a final volume of 1 ml. The cells were cultured in a CO₂ incubator at 37°C for 3 days. After completion of the cultivation, the total volume of cell suspension was recovered from each well and centrifuged (3,000 rpm, 1 min.) to recover the cells. The thus obtained cells were suspended in 100 μ l of PBS. Using 50 μ l of this suspension, specimens were prepared with a cell specimen preparing device, Cytospin3 (Shandon). After specimens were stained by the May-Grünwald-Giemsa staining method, 200 cells were observed for each specimen and the number of eosinophils was counted.

The results are shown in Fig. 58. Monoclonal antibodies KM1259 and KM1486 and humanized antibodies KM1399, KM7399, KM8399 and KM9399 were all found to have an activity to inhibit the eosinophil survival time prolongation by IL-5. However, such activity was not recognized in monoclonal antibody KM1257.

5. Detection of shIL-5R α with an Anti-hIL-5R α Antibodies

Anti-human IL-5R α monoclonal antibody KM1257 diluted with PBS to a concentration of 10 μ g/ml was dispensed into a 96-well EIA plate (Greiner) (50 μ l/well) and left at 4 °C overnight to allow the antibody to be adsorbed. After washing, 100 μ l/well of PBS containing 1% bovine serum albumin (BSA)(1% BSA-PBS) was added and reaction was performed at room temperature for 1 hour to block the remaining active groups. After discarding 1% BSA-PBS, the purified shlL-5R α obtained in subsection (9) of section 1 of Example 1 that had been diluted with 1% BSA-PBS to a concentration of 1000-0.1 ng/ml was added and reacted at 4 °C overnight. After washing with Tween-PBS, anti-human IL-5R α monoclonal antibody KM1259 labeled with biotin by conventional methods ["KOSO-KOTAI-HO" (Enzyme Antibody Method), Gakusai Kikaku Co., 1985] and diluted with 1% BSA-PBS to a concentration of 1 μ g/ml was added (50 μ l/well) and reacted at room temperature for 2 hours. After washing with Tween-PBS, avidin-labeled peroxidase (Nippon Reizo) diluted 4000 folds with 1% BSA-PBS was added (50 μ l/well) and reacted at room temperature for 1 hour. After washing with Tween-PBS, ABTS substrate solution [2,2-azinobis(3-ethylbenzothiazole-6-sulfonic acid)ammonium] was added to allow color development. Then, the absorbance at OD of 415 nm was measured (NJ2001; Japan Intermed).

The results are shown in Fig. 59. As a result, it has become clear that shIL-5R α can be measured by using anti-human IL-5R α monoclonal antibody KM1257 and biotin-labeled anti-human IL-5R α monoclonal antibody KM1259.

6. Detection of shIL-5R α by Western Blotting

The shIL-5R α described in subsection (9) of section 1 of Example 1 was thermally denatured in SDS-PAGE sample buffer containing 2-mercaptoethanol or SDS-PAGE sample buffer not containing 2-mercaptoethanol. The resultant mixture was electrophoresed on a commercial SDS-PAGE gradient gel (Atto), and then the protein was transferred to a PVDF membrane (Millipore). The PVDF membrane was soaked in PBS containing 10% BSA and left at 4 °C overnight for blocking. After completion of the blocking, the membrane was washed thoroughly with 0.05% Tween-containing PBS. Then, the membrane was soaked in a culture supernatant of the hybridoma obtained in section 5 of Example 1 at room temperature for 2 hours and washed thoroughly with 0.05% Tween-containing PBS. Further, the PVDF membrane was soaked at room temperature for 1 hour in a solution obtained by diluting peroxidase-labeled anti-mouse immunoglobulin antibody (Wako Pure Chemical Industries, Ltd.) with 1% BSA-PBS 1000 folds and then washed thoroughly with 0.05% Tween-containing PBS. After the washing solution was removed thoroughly, ECL reagent (Amersham) was

applied to the PVFD membrane and reacted for 1 minute. After removing the excessive reagent, the membrane was sandwiched between two plastic films and placed in an X-ray film sensitizing cassette to thereby sensitize the ECL film. Thus, the reactivity of the antibodies were confirmed.

The results are shown in Fig. 60. KM1257 exhibited reactivity, but KM1259 and KM1486 did not.

7. Immunoprecipitation of shIL-5R α

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An anti-mouse immunoglobulin antibody (DAKO) diluted with PBS 50 folds was dispensed into a 96-well EIA plastic plate (200 μ l/well) and left at 4°C overnight to allow the antibody to be adsorbed. After washing with PBS, 300 μ l/well of 1% BSA-PBS was added and left at room temperature for 1 hour to perform blocking. After washing with PBS, 200 μ l each of a culture supernatant of KM1257, KM1259 or KM1486 (they are anti-human IL-5R α monoclonal antibodies obtained in the preceding Examples) was added to each well and left at 4 °C overnight to allow the antibody to be adsorbed. After washing the plate, the shlL-5R α obtained in section 1 of Example 1 and diluted with PBS to a concentration of 10 μ g/ml was dispensed into each well in an amount of 50 μ l and reacted at 4°C overnight. After the plate was washed with 0.05% Tween-containing PBS, 5×2-mercaptoethanol-free SDS-PAGE sample buffer [0.31 M Tris (pH 6.8), 10% SDS, 50% glycerol] or 5×2-mercaptoethanol-containing SDS-PAGE sample buffer [0.31 M Tris (pH 6.8), 10% SDS, 50% glycerol, 25% 2-mercaptoethanol] was added (50 μ l/well) and left at room temperature for 2 hours while shaking. The reaction mixture was added to 200 μ l of PBS and heated on a heat block. Then, using a commercial SDS-PAGE gradient gel (Atto), 25 μ l of the reaction mixture was separated. After completion of the electrophoresis, the protein was transferred to a PVDF membrane (Millipore). The PVDF membrane was subjected to Western blotting analysis according to the method described in section 6 of Example 3 and using KM1257, to thereby detect shIL-5R α .

The results are shown in Fig. 61. It has become clear that all of KM1257, KM1259 and KM1486 immunoprecipitate shIL-5R α .

Industrial Applicability

According to the present invention, monoclonal antibodies KM1257, KM1259 and KM1486 are provided which specifically bind to human IL-5 receptor α chain that is believed to be specifically expressed on human eosinophils. Also, humanized antibodies KM1399, KM8399, KM7399 and KM9399 are provided which specifically bind to human IL-5 receptor α chain that is believed to be specifically expressed on human eosinophils and which can inhibit the biological activity of human IL-5. The antibodies of the present invention are useful for immunological detection of human eosinophils in immunocyte staining and diagnosis or treatment of allergic diseases caused by the inhibition of the biological activity of IL-5. It should be particularly noted that the humanized antibodies of the invention are lower in antigenicity than the monoclonal antibodies and expected to maintain their effect for a long period.

	Sequence Listing	
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	TOPOLOGY: linear	
10	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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15		
	SEQ ID NO: 2	
	SEQUENCE LENGTH: 32	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
25	SEQUENCE DESCRIPTION:	
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	SEQ ID NO: 3	
30	SEQUENCE LENGTH: 27	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single.	
35	TOPOLOGY: linear	
00	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CAGATATCTC ACTTCTCCCA CCTGTCA	27
40		
	SEQ ID NO: 4 .	
	SEQUENCE LENGTH: 88	
45	SEQUENCE TYPE: nucleic acid	
40	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
50	SEQUENCE DESCRIPTION:	
	AGCTTCCACC ATGGAGTTTG GGCTCAGCTG GCTTTTTCTT GTCCTTGTTT TCAAAGGTGT	60
	TCAGTGTGAC TTACTTCCTG ATGAAAAG	88

	SEQ ID NO: 5	
	SEQUENCE LENGTH: 84	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
10	SEQUENCE DESCRIPTION:	
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	GCTGAGCCCA AACTCCATGG TGGA	84
15		
	SEQ ID NO: 6	
	SEQUENCE LENGTH: 51	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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25	AGCTTCCACC ATGGCTACAG GCTCCCGGAC GTCCCTGCTC CTGGCTTTTG G	51
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30	SEQUENCE LENGTH: 58	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single-	
	TOPOLOGY: linear	
35	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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40	SEQ ID NO: 8.	
	SEQUENCE LENGTH: 64	
	SEQUENCE TYPE: nucleic acid	
45	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
50	CTTTTCATCA GGAAGTAAGT CGGCACTGCC CTCTTGAAGC CAGGGCAGGC AGAGCAGGCC	60
	AAAA	64
		-

	SEQ ID NO: 9	
	SEQUENCE LENGTH: 41	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
10	SEQUENCE DESCRIPTION:	
	GCCAGGAGCA GGGACGTCCG GGAGCCTGTA GCCATGGTGG A	41
15	SEQ ID NO: 10	
10	SEQUENCE LENGTH: 39	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
20	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	GGCAGCGGCG GTTCCGGTGA GCCCAAATCT TGTGACAAA	39
25		
	SEQ ID NO: 11	
	SEQUENCE LENGTH: 34	
30	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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40	SEQUENCE LENGTH: 34	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
45	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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50		
	SEQ ID NO: 13	
	SEQUENCE LENGTH: 39	

	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
5	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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10		
	SEQ ID NO: 14	
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	SEQUENCE TYPE: nucleic acid	
15	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
20	SEQUENCE DESCRIPTION:	
	CAAAGCTTCC ACCATGGCTA CAGGCTCCCG GACG	34
	SEQ ID NO: 15	
25	SEQUENCE LENGTH: 76	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: double	
	TOPOLOGY: linear	
30	MOLECULE TYPE: Other nucleic acid synthetic DNA	
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35	ACCCGGGCGC CATGCA	76
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	STRANDEDNESS: single	
	TOPOLOGY: linear	
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40	SEQUENCE DESCRIPTION:	
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50	SEQ ID NO: 17	
	SEQUENCE LENGTH: 40	
	SEQUENCE TYPE: nucleic acid	

	STRANDEDNESS: single												
	TOPOLOGY: linear												
5	MOLECULE TYPE: Other nucleic acid synthetic DNA												
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10	and TD 10 10												
10	SEQ ID NO: 18												
	SEQUENCE LENGTH: 21												
	SEQUENCE TYPE: nucleic acid												
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	MOLECULE TYPE: Other nucleic acid synthetic DNA												
	SEQUENCE DESCRIPTION:												
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20	ANTICOTACO GIGOLIGANO C												
	SEQ ID NO: 19												
	SEQUENCE LENGTH: 17												
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	TOPOLOGY: linear												
	MOLECULE TYPE: Other nucleic acid synthetic DNA												
30	SEQUENCE DESCRIPTION:												
	GGTGCAGCCA CCGTACG	17											
	SEQ ID NO: 20												
35	SEQUENCE LENGTH: 26												
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	STRANDEDNESS: single												
40	TOPOLOGY: linear												
	MOLECULE TYPE: Other nucleic acid synthetic DNA												
	SEQUENCE DESCRIPTION:												
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45													
	SEQ ID NO: 21												
	SEQUENCE LENGTH: 34												
50	SEQUENCE TYPE: nucleic acid												
	STRANDEDNESS: single												
	TOPOLOGY: linear												

	MOLECULE TYPE: Other nucleic acid synthetic DNA SEQUENCE DESCRIPTION:														
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5															
	SEQ ID NO: 22														
	SEQUENCE LENGTH: 421														
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	STRANDEDNESS: double														
	TOPOLOGY: linear														
15	MOLECULE TYPE: cDNA														
	FEATURE:														
	NAME/KEY: sig peptide														
	LOCATION: 157														
20	IDENTIFICATION METHOD: S														
	NAME/KEY: domain														
	LOCATION: 148162														
	IDENTIFICATION METHOD: S														
25	OTHER INFORMATION: CDR1														
	NAME/KEY: domain														
	LOCATION: 205255														
	IDENTIFICATION METHOD: S														
30	OTHER INFORMATION: CDR2														
	NAME/KEY: domain														
	LOCATION: 352387														
35	IDENTIFICATION METHOD: S														
	OTHER INFORMATION: CDR3														
	SEQUENCE DESCRIPTION:														
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40	Met Asn Phe Gly Leu Ser Leu Ile Phe Leu Ala Leu Ile Leu Lys Gly														
	-15 -10 -5														
	GTC CAA TGT GAG GTG CAG TTG GTG GAG TCT GGG GGA GAC TTA GTG AAG	96													
45	Val Gln Cys Glu Val Gln Leu Val Glu Ser Gly Gly Asp Leu Val Lys														
45	-1 1 5 10														
	CCT GGA GGG TCC CTG AAA CTC TCC TGT GCA GCC TCT GGA TTC ACT TTC	144													
	Pro Gly Gly Ser Leu Lys Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe														
50	15 20 25														
	AGT GAC TAT GGC ATG GCT TGG ATT CGC CAA ATT TCA GAC AAG AGG CCG	192													
	Ser Asp Tyr Gly Met Ala Trp Ile Arg Gln Ile Ser Asp Lys Arg Pro														

	30	30 35									40			45				
	GAG	TGG	GTC	GCA	GCC	ATT	AGC	AGT	GGT	GGT	AGT	TAC	ATC	CAC	TTT	CCA	240	
5	Glu	Trp	Val	Ala	Ala	Ile	Ser	Ser	Gly	Gly	Ser	Tyr	Ile	His	Phe	Pro		
					50					55					60			
	GAC	AGT	TTG	AAG	GGG	CGA	TTC	ACC	GTC	TCC	AGA	GAC	AAT	GCC	AAG	AAC	288	
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10				65					70					75				
	ACC	CTG	TAC	CTG	GAA	ATG	AGC	GGT	CTG	AAG	TCT	GAG	GAC	ACA	GCT	ATG	336	
	Thr	Leu	Tyr	Leu	Glu	Met	Ser	Gly	Leu	Lys	Ser	Glu	Asp	Thr	Ala	Met		
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	Tyr	Tyr	Cys	Ala	Arg	Arg	Gly	Phe	Tyr	Gly	Asn	Tyr	Arg	Ala	Met	Asp		
		95					100					105						
20	TAC	TGG	GGT	CAA	GGA	ACC	TCA	GTC	ACC	GTC	TCC	TCA	G				421	
	Tyr	Trp	Gly	Gln	Gly	Thr	Ser	Val	Thr	Val	Ser	Ser						
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05																		
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30	STRANDEDNESS: double																	
	TOPOLOGY: linear																	
	MOLECULE TYPE: cDNA.																	
	FEATURE:																	
35	NAME/KEY: sig peptide																	
	LOCATION: 160																	
	IDEN	TIFI	CATI	ON M	ETHO	D: S										•		
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40	LOCA'	TION	: 13	01	74													
	IDENTIFICATION METHOD: S																	
	OTHE	R IN	FORM	ATIO	N: C	DR1												
45	NAME	/KEY	: do	main														
	LOCA	rion	: 22	02	40										•			
	IDEN	rifi	CATI	ON M	ETHO	D: S												
	OTHER	RIN	FORM	ATIO	N: C	DR2												
50	NAME	/KEY	: do	main														
	LOCAT	rion	: 33	73	63													
	IDENT	rifi	CATI	M NC	ETHO	D: S												

	OTH	ER I	NFOR	MATI	ON:	CDR3											
	SEQ	UENC	E DE	SCRI	PTIO	N:											
5	ATG	GAG	AAA	GAC	ACA	CTC	CTG	CTA	TGG	GTC	CTG	CTT	CTC	TGG	GTT	CCA	48
	Met	Glu	Lys	Asp	Thr	Leu	Leu	Leu	Trp	Val	Leu	Leu	Leu	Trp	Val	Pro	
	-20			•		-15					-10					-5	
	GGT	TCC	AGA	AGT	GAC	ATT	GTG	CTG	ACC	CAA	TCT	CCA	GCT	TCT	TTG	GCT	96
10	Gly	Ser	Arg	Ser	Asp	Ile	Val	Leu	Thr	Gln	Ser	Pro	Ala	Ser	Leu	Ala	
				-1	1		•		5					10			
	GTG	TCT	CTA	GGG	CAG	AGG	GCC	ACC	ATC	TCC	TGC	AGA	GCC	AAC	GAA	AGT	144
15	Val	Ser	Leu	Gly	Gln	Arg	Ala	Thr	Ile	Ser	Cys	Arg	Ala	Asn	Glu	Ser	
			15					20					25				
	GTT	GAT	CAT	AAT	GGC	GTC	AAT	TTT	ATG	AAC	TGG	TTC	CAA	CAG	AAA	CCA	192
	Val	Asp	His	Asn	Gly	Val	Asn	Phe	Met	Asn	Trp	Phe	Gln	Gln	Lys	Pro	
20		30					35					40					
	GGA	CAG	TCA	CCC	AAG	CTC	CTC	ATC	TAT	GCT	GCA	TCC	AAC	CAA	GGA	TCC	240
	Gly	Gln	Ser	Pro	Lys	Leu	Leu	Ile	Tyr	Ala	Ala	Ser	Asn	Gln	Gly	Ser	
	45					50					55					60	
25	GGC	GTC	CCT	GCC	AGG	TTT	AGT	GGC	AGT	GGG	TCT	GGG	ACA	GAC	TTC	AGT	288
	Gly	Val	Pro	Ala	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Ser	
					65					70					75		
30	CTC	AAC	ATC	CAT	CCT	ATG	GAG	GAG	GAT	GAT	GCT	GCA	ATG	TAT	TTC	TGT	336
	Leu	Asn	Ile	His	Pro	Met	Glu	Glu	Asp	Asp	Ala	Ala	Met	Tyr	Phe	Cys	
				80					85					90			
	CAG	CAA	AGT	AAG	GAT	GTT	CCG	TGG	ACG	TTC	GGT	GGA	GGC	ACC	AGG	TTG	384
35	Gln	Gln	Ser	Lys	Asp	Val	Pro	Trp	Thr	Phe	Gly	Gly	Gly	Thr	Arg	Leu	
			95					100					105				
	GAA	ATC	AAA	С													394
10	Glu	Ile	Lys														
40		110	111	•													
	SEQ	ID N	10: 2	24													
45	SEQU	ENCE	LEN	IGTH:	421	-											
	SEQU	ENCE	TYF	E: r	ucle	eic a	cid										
	STRA	NDED	NESS	G: do	uble	:											
	TOPO	LOGY	: li	.near	•												
	MOLE	CULE	TYP	E: c	:DNA												
	FEAT	URE:															
	NAME	/KEY	: si	g pe	ptid	e											

	LOC	OITA	N: 1	57													
	IDE	NTIF	'ICAT	'ION	метн	OD:	S										
5	NAM	E/KE	Y: d	omai.	.n												
	LOC	ATIO	N: 1	48	162												
	IDE	NTIF	ICAT	ION	метн	OD:	S										
	ОТН	ER I	NFOR	MATI	ON:	CDR1											
10	NAM	E/KE	Y: d	omai	n												
	LOC	ATIO	N: 2	05	255												
	IDE	NTIF	ICAT	ION	METH	OD:	S										
	ОТН	ER I	NFOR	MATI	ON:	CDR2											
15	NAM	E/KE	Y: d	omai	n												
	LOC	ATIO	N: 3	52	387												
	IDE	NTIF	ICAT	ION	METH	OD:	s										
20	OTH	ER I	NFOR	MATI	ON:	CDR3											
	SEQ	UENC	E DE	SCRI	PTIO	N:											
	ATG	GAA	TGG	AGT	TGG	ATA	TTT	CTC	TTT	CTC	CTG	TCA	GGA	ACT	GCA	GGT	48
	Met	Glu	Trp	Ser	Trp	Ile	Phe	Leu	Phe	Leu	Leu	Ser	Gly	Thr	Ala	Gly	
25					-15					-10					-5		
	GTC	CAC	TCT	GAG	GTC	CAG	CTG	CAA	CAG	TCT	GGA	CCT	GAG	CTG	GTA	AAG	96
	Val	His	Ser	Glu	Val	Gln	Leu	Gln	Gln	Ser	Gly	Pro	Glu	Leu	Val	Lys	
20			-1	1				5					10				
30	CCT	GGG	GCT	TCA	GTG	AAG	ATG	TCC	TGC	AAG	GCT	TCT	GGA	TAC	ACA	TTC	144
	Pro	Gly	Ala	Ser	Val	Lys	Met	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Thr	Phe	
		15					20					25					
35	ACT	AGT	TAT	GTT	ATT	CAC	TGG	GTG	AAA	CAG	AGG	CCT	GGT	CAG	GGC	CTT	192
	Thr	Ser	Tyr	Val	Ile	His	Trp	Val	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	
	30					35					40					45	
	GCG	TGG	ATT	GGA	TAT	ATT	AAT	CCT	TAC	AAT	GAT	GGG	ACT	AAG	TAC	AAT	240
40	Ala	Trp	Ile	Gly	Tyr	Ile	Asn	Pro	Tyr	Asn	Asp	Gly	Thr	Lys	Tyr	Asn	
					50					55					60		
	GAG	AGG	TTC	AAA	GGC	AAG	GCC	ACA	CTG	ACT	TCA	GAC	AGA	TCC	TCC	AGC	288
45	Glu	Arg	Phe	_	Gly	Lys	Ala	Thr	Leu	Thr	Ser	Asp	Arg	Ser	Ser	Ser	
				65					70					75			
												GAG					336
	Thr	Val	-	Met	Glu	Leu	Ser		Leu	Thr	Ser	Glu		Ser	Ala	Val	
50			80			_		85					90				
												GGT					384
	Tyr	Leu	Cys	Gly	Arg	Glu	Gly	Ile	Arg	Tyr	Tyr	Gly	Leu	Leu	Gly	Asp	

	95 100 105														
	TAC TGG GGC CAA GGC ACC ACT CTC ACA GTC TCC TCA G	21													
5	Tyr Trp Gly Gln Gly Thr Thr Leu Thr Val Ser Ser														
	110 115 120 121														
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10	SEQUENCE LENGTH: 382														
	SEQUENCE TYPE: nucleic acid														
	STRANDEDNESS: double														
15	TOPOLOGY: linear														
10	MOLECULE TYPE: cDNA														
	FEATURE:														
	NAME/KEY: sig peptide														
20	LOCATION: 160														
	IDENTIFICATION METHOD: S														
	NAME/KEY: domain														
	LOCATION: 130162														
25	IDENTIFICATION METHOD: S														
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	NAME/KEY: domain														
20	LOCATION: 208228														
30	IDENTIFICATION METHOD: S														
	OTHER INFORMATION: CDR2														
	NAME/KEY: domain														
35	LOCATION: 325351														
	IDENTIFICATION METHOD: S														
	OTHER INFORMATION: CDR3														
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40	ATG ATG TCC TCT GCT CAG TTC CTT GGT CTC CTG TTG CTC TGT TTT CAA	48													
	Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln														
	-20 -15 -10 -5														
45	GAT ATC AGA TGT GAT ATC CAG ATG ACA CAG GCT ACA TCC TCC CTG TCT	96													
40	Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ala Thr Ser Ser Leu Ser														
	-1 1 5 10														
	GCC TCT CTG GGA GAC AGA GTC ACC ATC GGT TGC GGG ACA AGT GAG GAC	14													
50	Ala Ser Leu Gly Asp Arg Val Thr Ile Gly Cys Gly Thr Ser Glu Asp														
	15 20 25														
	ATT ATC AAT TAT TTA AAC TGG TAT CGG AAG AAA CCA GAT GGA ACT GTT 19)2													

	Ile 1		sn Tyr	Leu	Asn		туr	Arg	Lys	Lys		Asp	Gly	Thr	Val	
		30				35					40					
5			rg atc													240
		Leu Le	eu Ile	Tyr		Thr	Ser	Arg	Leu		Ser	Gly	Val	Pro		
	45				50					55					60	
10	AGG I	TC AC	ST GGC	AGC	GGG	TCT	GGA	ACA	GAT	TAT	TCT	CTC	ACC	ATT	AGT	288
10	Arg F	Phe Se	er Gly	Ser	Gly	Ser	Gly	Thr	Asp	Tyr	Ser	Leu	Thr	Ile	Ser	
				65					70					75		
			AG CAA													336
15	Asp L	Leu Gl	u Gln	Glu	Asp	Ile	Ala	Thr	Tyr	Phe	Cys	Gln	Gln	Gly	Tyr	
			80					85					90			
	ACG C	CTT CC	G TAC	ACG	GTC	GGA	GGG	GGG	ACC	AAG	TTG	GAA	ATA	AAA	С	382
	Thr L		o Tyr	Thr	Val	Gly	Gly	Gly	Thr	Lys	Leu		Ile	Lys		
20		9	5				100					105		107		
	SEQ ID NO: 26															
25			ENGTH													
25	SEQUE	NCE T	YPE: r	nucle	eic a	cid										
	STRAN	IDEDNE	SS: do	ouble	<u> </u>											
	TOPOLOGY: linear															
30	MOLECULE TYPE: cDNA															
	FEATU	RE:														
	NAME/KEY: sig peptide															
	LOCATION: 157															
35	IDENTIFICATION METHOD: S															
	NAME/KEY: domain															
	LOCATION: 148162															
	IDENTIFICATION METHOD: S															
40	OTHER INFORMATION: CDR1															
	NAME/KEY: domain															
	LOCAT	ION:	2052	55												
45	IDENT	IFICA	TION M	ETHO	D: S											
	OTHER	INFO	RMATIO	N: C	DR2											
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	LOCATI	ION:	3523	78												
50	IDENT	IFICA	TION M	ETHO	D: S											
	OTHER	INFO	RMATIO	N: C	DR3											
	SEQUE	NCE DI	ESCRIP	TION	:											

	ATG	AAA	TGC	AGC	TGG	GTT	ATC	TTC	TTC	CTG	ATG	GCA	GTG	GTT	ACA	GGG	48
						Val											
5					-15					-10					-5		
	GTC	AAT	TCA	GAG	GTT	CAG	CTG	CAG	CAG	TCT	GGG	GCA	GAG	CTT	GTG	AAG	96
	Val	Asn	Ser	Glu	Val	Gln	Leu	Gln	Gln	Ser	Gly	Ala	Glu	Leu	Val	Lys	
			-1	1				5					10				
10						AAC											144
	Pro	Gly	Ala	Ser	Val	Asn	Leu	Ser	Cys	Thr	Ala	Ser	Gly	Phe	Asn	Ile	
		15					20					25					
15						CAC											192
,,,	Lys	Asp	Thr	Tyr	Met	His	Trp	Val	Lys	Gln		Pro	Glu	Gln	Gly		
	30					35					40					45	
						ATT											240
20	Glu	Trp	Ile	Gly		Ile	Asp	Pro	Ala		Gly	Asn	Thr	Lys		Asp	
					50					55	CC.	C N C	3.03	mcc.	60	220	288
						AAG											200
05	Pro	Lys	Pne		Ala	Lys	Ala	THE	70	Ald	Ald	wsp	1111	75	Ser	ASII	
25			ma a	65 CmC	CAC	CTC	NCC	NCC.		አሮአ	ጥርጥ	GAG	GAC		GCC	GTC	336
						Leu											330
	1111	Ald	80	Ten	GIII	Dea	Jei	85	БСи			014	90				
30	тΔт	TAC		ACT	GGT	GGA	СТА		CTA	CGG	TTC	TTT		TAT	TGG	GGC	384
						Gly											
	-1-	95	-1-		1		100	,		-		105	-	_	_	_	
	CAA		ACC	ACT	CTC	ACA	GTC	TCC	TCA	G							412
35						Thr											
	110					115			118							•	
40	SEQ	ID N	10: 2	27													
	SEQU	SEQ ID NO: 27 SEQUENCE LENGTH: 331															
	SEQU	JENCE	TY	PE: r	nucle	eic a	acid										
	STRA	ANDEI	ONESS	3: do	ouble	2											
45	TOPO	DLOGY	(: li	inear	:												
	MOLE	CULE	TY	PE: 0	DNA												
	FEAT	CURE:	;														
50	NAME	E/KEY	(: do	omair	1												
50	LOCA	OITA	1: 79	910	8												
	IDEN	TIF	CAT	ON N	(ETH	DD: S	3										

	OTH	ER I	NFOR	MATI	ON:	CDR1											
	NAM	E/KE	Y: đ	omai	n												
5	LOC	ATIO	N: 1	54	174												
	IDE	NTIF	ICAT	ION	METH	OD:	s										
	OTH	ER I	NFOR	MAT'I	ON:	CDR2											
	NAM	E/KE	Y: d	omai	n												
10	LOC	ATIO	N: 2	71	300												
	IDE	NTIF	ICAT	ION	METH	OD:	s										
	ОТН	ER I	NFOR	MATI	ON:	CDR3											
	SEQ	UENC	E DE	SCRI	PTIO	N:											
15	TCC	AGA	GGA	CAA	ATT	GTT	CTC	ACC	CAG	TCT	CCA	GCA	ATC	ATG	TCT	GCA	48
	Ser	Arg	Gly	Gln	Ile	Val	Leu	Thr	Gln	Ser	Pro	Ala	Ile	Met	Ser	Ala	
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20	TCT	CCA	GGG	GAG	AAG	GTC	ACC	ATG	ACC	TGC	AGT	GCC	AGT	TCA	AGT	GTA	96
	Ser	Pro	Gly	Glu	Lys	Val	Thr	Met	Thr	Cys	Ser	Ala	Ser	Ser	Ser	Val	
		15					20					25					
	AGT	TAC	ATG	CAC	TGG	TAC	CAG	CAG	AAG	TCA	GGC	ACC	TCC	CCC	AAA	AGA	144
25	Ser	Tyr	Met	His	Trp	Tyr	Gln	Gln	Lys	Ser	Gly	Thr	Ser	Pro	Lys	Arg	
	30					35					40					45	
	TGG	ATT	TAT	GAC	ACA	TCC	AAA	CTG	GCT	TCT	GGA	GTC	CCT	GCT	CGC	TTC	192
30	Trp	Ile	Tyr	Asp	Thr	Ser	Lys	Leu	Ala	Ser	Gly	Val	Pro	Ala	Arg	Phe	
50					50					55					60		
	AGT	GGC	AGT	GGG	TCT	,GGG	ACC	TCT	TAC	TCT	CTC	ACA	ATC	AGC	AGC	ATG	240
	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Ser	Tyr	Ser	Leu	Thr	Ile	Ser	Ser	Met	
35				65					70					75			
	GAG	GCT	GAA	GAT	GCT	GCC	ACT	TAT	TAC	TGC	CAG	CAG	TGG	AGT	AGT	AAC	288
	Glu	Ala	Glu	Asp	Ala	Ala	Thr	Tyr	Tyr	Cys	Gln	Gln	Trp	Ser	Ser	Asn	
			80					85					90				
40	CCA	CCC	ATC	ACG	TTC	GGA	GGG	GGG	ACC	AAG	CTG	GAA	ATA	AAA	С		331
	Pro	Pro	Ile	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys			
		95					100					105		107			
45																	
	SEQ	ID N	10: 2	28													
	SEQU	JENCE	LEN	IGTH :	5												
	SEQU	JENCE	TYF	E: a	mino	aci	.d										
50	STRA	NDED	NESS	: si	ngle	•											
	TOPO	LOGY	: li	.near	•												
	MOT.F	CULE	TYP	e: r	epti	đe											

	SEQUENCE DESCRIPTION:
	Asp Tyr Gly Met Ala
5	
	SEQ ID NO: 29
	SEQUENCE LENGTH: 17
	SEQUENCE TYPE: amino acid
10	STRANDEDNESS: single
	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
15	Ala Ile Ser Ser Gly Gly Ser Tyr Ile His Phe Pro Asp Ser Leu Lys Gly
	SEQ ID NO: 30
20	SEQUENCE LENGTH: 12
	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
25	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
	Arg Gly Phe Tyr Gly Asn Tyr Arg Ala Met Asp Tyr
30	SEQ ID NO: 31
	SEQUENCE LENGTH: 15.
	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
35	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
40	Arg Ala Asn Glu Ser Val Asp His Asn Gly Val Asn Phe Met Asn
	SEQ ID NO: 32
	SEQUENCE LENGTH: 7
45	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
50	MOLECULE TYPE: peptide
•	SEQUENCE DESCRIPTION:
	Ala Ala Ser Asn Gln Gly Ser

	SEQ ID NO: 33
	SEQUENCE LENGTH: 9
5	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
	MOLECULE TYPE: peptide
10	SEQUENCE DESCRIPTION:
	Gln Gln Ser Lys Asp Val Pro Trp Thr
15	SEQ ID NO: 34
	SEQUENCE LENGTH: 5
	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
20	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
25	Ser Tyr Val Ile His
	SEQ ID NO: 35
	SEQUENCE LENGTH: 17
30	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single 1
	TOPOLOGY: linear
35	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
	Tyr Ile Asn Pro Tyr Asn Asp Gly Thr Lys Tyr Asn Glu Arg Phe Lys Gly
40	SEQ ID NO: 36
	SEQUENCE LENGTH: 12
	SEQUENCE TYPE: amino acid
45	STRANDEDNESS: single
	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
50	Glu Gly Ile Arg Tyr Tyr Gly Leu Leu Gly Asp Tyr

64

SEQ ID NO: 37

	SEQUENCE LENGTH: 11
	SEQUENCE TYPE: amino acid
5	STRANDEDNESS: single
	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
10	Gly Thr Ser Glu Asp Ile Ile Asn Tyr Leu Asn
	SEQ ID NO: 38
15	SEQUENCE LENGTH: 7
	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
20	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
	His Thr Ser Arg Leu Gln Ser
0.5	
25	SEQ ID NO: 39
	SEQUENCE LENGTH: 9
	SEQUENCE TYPE: amino acid
30	STRANDEDNESS: single
	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
35	Gln Gln Gly Tyr Thr Leu Pro Tyr Thr
	· ·
	SEQ ID NO: 40
	SEQUENCE LENGTH: 5
40	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
45	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
	Asp Thr Tyr Met His
50	SEQ ID NO: 41
	SEQUENCE LENGTH: 17
	SEQUENCE TYPE: amino acid

	STRANDEDNESS: single
	TOPOLOGY: linear
5	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
	Arg Ile Asp Pro Ala Asn Gly Asn Thr Lys Ser Asp Pro Lys Phe Gln Ala
10	SEQ ID NO: 42
	SEQUENCE LENGTH: 9
	SEQUENCE TYPE: amino acid
15	STRANDEDNESS: single
15	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
20	Gly Leu Arg Leu Arg Phe Phe Asp Tyr
	SEQ ID NO: 43
	SEQUENCE LENGTH: 10
25	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear
30	MOLECULE TYPE: peptide
00	SEQUENCE DESCRIPTION:
	Ser Ala Ser Ser Ser Val Ser Tyr Met His
35	SEQ ID NO: 44
	SEQUENCE LENGTH: 7
	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
40	TOPOLOGY: linear
	MOLECULE TYPE: peptide
	SEQUENCE DESCRIPTION:
45	Asp Thr Ser Lys Leu Ala Ser
	SEQ ID NO: 45
	SEQUENCE LENGTH: 10
50	SEQUENCE TYPE: amino acid
	STRANDEDNESS: single
	TOPOLOGY: linear

	MOLECULE TYPE: peptide	
	SEQUENCE DESCRIPTION:	
5	Gln Gln Trp Ser Ser Asn Pro Pro Ile Thr	
	SEQ ID NO: 46	
	SEQUENCE LENGTH: 39	
10	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
15	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	GCACCACTCT CACAGTCTCC TCAGCCAGTA CTAAGGGCC	39
20	SEQ ID NO: 47	
	SEQUENCE LENGTH: 31	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
25	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
30	CTTAGTACTG GCTGAGGAGA CTGTGAGAGT G	31
	SEQ ID NO: 48	
	SEQUENCE LENGTH: 20	
35	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
40	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	20
	GACCAAGTTG GAAATAAAAC	20
45	SEQ ID NO: 49	
45	SEQUENCE LENGTH: 21	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
50	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	

	GTACGTTTTA TTTCCAACTT G	21
5	SEQ ID NO: 50	
	SEQUENCE LENGTH: 97	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
10	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
15	CAGGAAACAG CTATGACGCG GCCGCCACCA TGGAATGGAG TTGGATATTT CTCTTTCTCC	60
15	TGTCAGGAAC TGCAGGTGTC CACTCTGAGG TCCAGCT	97
	SEQ ID NO: 51	
20	SEQUENCE LENGTH: 96	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
25	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	GAATGTGTAT CCAGAAGCCT TGCAGGAAAC CTTCACTGAA GCCCCAGGCT TCTTCACCTC	60
	AGCTCCAGAC TGCACCAGCT GGACCTCAGA GTGGAC	96
30		
	SEQ ID NO: 52	
	SEQUENCE LENGTH: 96	
35	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
40	SEQUENCE DESCRIPTION:	
	AGGCTTCTGG ATACACATTC ACTAGTTATG TTATTCACTG GGTGCGACAG GCCCCTGGTC	60
	AGGGCCTTGA GTGGATGGGA TATATTAATC CTTACA	96
45	270 TD 170 - 53	
	SEQ ID NO: 53	
	SEQUENCE LENGTH: 98	
	SEQUENCE TYPE: nucleic acid	
50	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	

	SEQUENCE DESCRIPTION:	
	TGTAGGCTGT GCTCGTGGAC GTGTCTGCAG TGATTGTGAC TCTGCCTTTG AACCTCTCAT	60
5	TGTACTTAGT CCCATCATTG TAAGGATTAA TATATCCC	96
	SEQ ID NO: 54	
	SEQUENCE LENGTH: 96	
10	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
15	MOLECULE TYPE: Other nucleic acid synthetic DNA	
15	SEQUENCE DESCRIPTION:	
	GTCCACGAGC ACAGCCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA	60
	TTACTGTGCG AGAGAAGGAA TTAGGTACTA TGGTCT	96
20		
	SEQ ID NO: 55	
	SEQUENCE LENGTH: 100	
	SEQUENCE TYPE: nucleic acid	
25	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
30	SEQUENCE DESCRIPTION:	
30	GTTTTCCCAG TCACGACGGG CCCTTGGTGG AGGCTGAGGA GACTGTGACC AGGGTGCCTT	60
	GGCCCCAGTA GTCTCCCAGT AGACCATAGT ACCTAATTCC	100
35	SEQ ID NO: 56	
	SEQUENCE LENGTH: 421	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: double	
40	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	FEATURE:	
45	NAME/KEY: sig peptide	
	LOCATION: 157	
	IDENTIFICATION METHOD: S	
	NAME/KEY: domain	
50	LOCATION: 148162	
	IDENTIFICATION METHOD: S	
	OTHER INFORMATION: CDR1	

	NAM	ie/ke	Y: c	lomai	in												
	LOC	CATIC	N: 2	205	255												
5	IDE	NTIF	ICAT	CION	METH	OD:	s										
	OTH	ER I	NFOF	TAMS	ON:	CDR 2	?										
	NAM	E/KE	Y: d	lomai	.n												
	LOC	ATIC	N: 3	52	378												
10	IDE	NTIF	'ICAT	NOI	метн	OD:	S										
	OTH	ER I	NFOR	MATI	ON:	CDR3											
	SEQ	UENC	E DE	SCRI	PTIO	N:											
	ATG	GAA	TGG	AGT	TGG	ATA	TTT	CTC	TTT	CTC	CTG	TCA	GGA	ACT	GCA	GGT	48
15	Met	Glu	Trp	Ser	Trp	Ile	Phe	Leu	Phe	Leu	Leu	Ser	Gly	Thr	Ala	Gly	
					-15					-10					-5		
	GTC	CAC	TCT	GAG	GTC	CAG	CTG	GTG	CAG	TCT	GGA	GCI	GAG	GTG	AAG	AAG	96
20	Val	His	Ser	Glu	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys	
			-1	1				5					10				
	CCT	GGG	GCT	TCA	GTG	AAG	GTT	TCC	TGC	AAG	GCT	TCT	GGA	TAC	ACA	TTC	144
	Pro	Gly	Ala	Ser	Val	Lys	Val	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Thr	Phe	
25		15					20					25					
	ACT	AGT	TAT	GTT	ATT	CAC	TGG	GTG	CGA	CAG	GCC	CCT	GGT	CAG	GGC	CTT	192
	Thr	Ser	Tyr	Val	Ile	His	Trp	Val	Arg	Gln	Ala	Pro	Gly	Gln	Gly	Leu	
	30					35					40					45	
30	GAG	TGG	ATG	GGA	TAT	ATT	AAT	CCT	TAC	AAT	GAT	GGG	ACT	AAG	TAC	AAT	240
	Glu	Trp	Met	Gly	Tyr	I·le	Asn	Pro	Tyr	Asn	Asp	Gly	Thr	Lys	Tyr	Asn	
					50					55					60		
35	GAG	AGG	TTC	AAA	GGC	AGA	GTC	ACA	ATC	ACT	GCA	GAC	ACG	TCC	ACG	AGC	288
	Glu	Arg	Phe	Lys	Gly	Arg	Val	Thr	Ile	Thr	Ala	Asp	Thr	Ser	Thr	Ser	
				65					70					75			
	ACA	GCC	TAC	ATG	GAG	CTC	AGT	TCG	CTG	AGA	TCT	GAG	GAC	ACG	GCG	GTG	336
40	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	Leu	Arg	Ser	Glu	Asp	Thr	Ala	Val	
			80					85					90				
	TAT	TAC	TGT	GCG	AGA	GAA	GGA	ATT	AGG	TAC	TAT	GGT	CTA	CTG	GGA	GAC	384
	Tyr	Tyr	Cys	Ala	Arg	Glu	Gly	Ile	Arg	Tyr	Tyr	Gly	Leu	Leu	Gly	Asp	
45		95					100					105					
	TAC	TGG	GGC	CAA	GGC	ACC	CTG	GTC	ACA	GTC	TCC	TCA	G				421
	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Val	Thr	Val	Ser	Ser					
50	110					115					120	121					

SEQ ID NO: 57

	SEQUENCE LENGTH: 87	
	SEQUENCE TYPE: nucleic acid	
5	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
10	CAGGAAACAG CTATGACGAA TTCCACCATG ATGTCCTCTG CTCAGTTCCT TGGTCTCCTG	60
	TTGCTCTGTT TTCAAGACAT CAGATGT	87
15	SEQ ID NO: 58	
	SEQUENCE LENGTH: 83	
	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
20	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
25	GATGGTGACT CTGTCTCCTA CAGAGGCAGA CAGGGAGGAT GGAGACTGTG TCATCTGGAT	60
20	ATCACATCTG ATGTCTTGAA AAC	83
	SEQ ID NO: 59	
	SEQUENCE LENGTH: 92	
30	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single.	
	TOPOLOGY: linear	
35	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	TAGGAGACAG AGTCACCATC ACTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60
	GGTATCAACA GAAACCAGGG AAAGCCCCTA AG	92
40		
	SEQ ID NO: 60	
	SEQUENCE LENGTH: 90	
45	SEQUENCE TYPE: nucleic acid	
45	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
50	SEQUENCE DESCRIPTION:	
	TTCCAGACCC GCTGCCACTG AACCTTGATG GGACTCCTGA CTGTAATCTT GATGTGTGGT	60
	AGATCAGGAG CTTAGGGGCT TTCCCTGGTT	90
	,	

	SEQ ID NO: 61	
	SEQUENCE LENGTH: 88	
5	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
10	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTAGTCTGC AACCTGAAGA	60
	TTTTGCCACT TACTACTGCC AACAGGGT	88
15		
,,,	SEQ ID NO: 62	
	SEQUENCE LENGTH: 91	
	SEQUENCE TYPE: nucleic acid	
20	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
05	SEQUENCE DESCRIPTION:	
25	GTTTTCCCAG TCACGACCGT ACGTTTTATT TCCACCTTGG TCCCTTGGCC GAACGTGTAC	60
	GGAAGCGTAT AACCCTGTTG GCAGTAGTAA G	91
30	SEQ ID NO: 63	
	SEQUENCE LENGTH: 382.	
	SEQUENCE TYPE: nucleic acid	
35	STRANDEDNESS: double	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
40	FEATURE:	
	NAME/KEY: sig. peptide	
	LOCATION: 160	
45	IDENTIFICATION METHOD: S	
	NAME/KEY: domain	
	LOCATION: 130162	
	IDENTIFICATION METHOD: S	
50	OTHER INFORMATION: CDR1	
	NAME/KEY: domain	
	LOCATION: 208228	
	IDENTIFICATION METHOD: S	

	OTH	ER I	NFOR	MATI	:NC	CDR2											
	NAMI	E\KE	Y: do	omai	n												
5	LOC	ATIO	N: 3	25	351												
	IDE	NTIF	ICAT:	ION (METH	DD:	S										
	OTH	ER I	NFOR	MATIO	: NC	CDR3											
	SEQ	JENC	E DES	SCRI	PTIO	N:											
10	ATG	ATG	TCC	TCT	GCT	CAG	TTC	CTT	GGT	CTC	CTG	TTG	CTC	TGT	TTT	CAA	48
	Met	Met	Ser	Ser	Ala	Gln	Phe	Leu	Gly	Leu	Leu	Leu	Leu	Cys	Phe	Gln	
	-20					-15					-10					-5	
	GAT	ATC	AGA	TGT	GAT	ATC	CAG	ATG	ACA	CAG	TCT	CCA	TCC	TCC	CTG	TCT	96
15	Asp	Ile	Arg	Cys	Asp	Ile	Gln	Met	Thr	Gln	Ser	Pro	Ser	Ser	Leu	Ser	
				-1	1				5					10			
	GCC	TCT	GTA	GGA	GAC	AGA	GTC	ACC	ATC	ACT	TGC	GGG	ACA	AGT	GAG	GAC	144
20	Ala	Ser	Val	Gly	Asp	Arg	Val	Thr	Ile	Thr	Cys	Gly	Thr	Ser	Glu	Asp	
			15					20					25				
	ATT	ATC	AAT	TAT	TTA	AAC	TGG	TAT	CAA	CAG	AAA	CCA	GGG	AAA	GCC	CCT	192
	Ile	Ile	Asn	Tyr	Leu	Asn	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Lys	Ala	Pro	
25		30					35					40					
	AAG	CTC	CTG	ATC	TAC	CAC	ACA	TCA	AGA	TTA	CAG	TCA	GGA	GTC	CCA	TCA	240
	Lys	Leu	Leu	Ile	Tyr	His	Thr	Ser	Arg	Leu	Gln	Ser	Gly	Val	Pro	Ser	
	45					50					55					60	
30	AGG	TTC	AGT	GGC	AGC	GGG	TCT	GGA	ACA	GAT	TTC	ACT	CTC	ACC	ATT	AGT	288
	Arg	Phe	Ser	G1y	Ser	-Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Ser	
					65					70					75		
35	AGT	CTG	CAA	CCT	GAA	GAT	TTT	GCC	ACT	TAC	TAC	TGC	CAA	CAG	GGT	TAT	336
	Ser	Leu	Gln	Pro	Glu	Asp	Phe	Ala	Thr	Tyr	Tyr	Cys	Gln	Gln	Gly	Tyr	
				80					85					90			
	ACG	CTT	CCG	TAC	ACG	TTC	GGC	CAA	GGG	ACC	AAG	GTG	GAA	ATA	AAA	С	382
40	Thr	Leu	Pro	Tyr	Thr	Phe	Gly	Gln	Gly	Thr	Lys	Val	Glu	Ile	Lys		
			95					100					105		107		
	SEQ	ID N	10: 6	54													
45	SEQU	JENCE	E LEN	GTH:	96												
	SEQU	JENCE	TYE	PE: r	nucle	eic a	acid										
			NESS														
50			: li		-												
	MOLE	CULI	TYE	E: C	ther	nuc	cleid	aci	id s	ynth	etic	DNA	1				
			DES							-							

SEQ ID NO: 65 SEQUENCE LENGTH: 421 SEQUENCE TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S OTHER INFORMATION: CDR1	96														
SEQ ID NO: 65 SEQUENCE LENGTH: 421 SEQUENCE TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
SEQUENCE LENGTH: 421 SEQUENCE TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
SEQUENCE TYPE: nucleic acid STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
TOPOLOGY: linear MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
FEATURE: NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
NAME/KEY: sig peptide LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
LOCATION: 157 IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
IDENTIFICATION METHOD: S NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
NAME/KEY: domain LOCATION: 148162 IDENTIFICATION METHOD: S															
LOCATION: 148162 IDENTIFICATION METHOD: S															
IDENTIFICATION METHOD: S															
NAME/KEY: domain															
LOCATION: 205255															
IDENTIFICATION METHOD: S															
OTHER INFORMATION: CDR2															
30 NAME/KEY: domain															
LOCATION: 352378 -	LOCATION: 352378 -														
IDENTIFICATION METHOD: S															
OTHER INFORMATION: CDR3															
SEQUENCE DESCRIPTION:															
ATG GAA TGG AGT TGG ATA TTT CTC TTT CTC CTG TCA GGA ACT GCA GGT	48														
Met Glu Trp Ser Trp Ile Phe Leu Phe Leu Leu Ser Gly Thr Ala Gly															
<i>40</i> · −15 −10 −5															
GTC CAC TCT GAG GTC CAG CTG GTG CAG TCT GGA GCT GAG GTG AAG AAG	96														
Val His Ser Glu Val Gln Leu Val Gln Ser Gly Ala Glu Val Lys Lys															
-1 1 5 10															
45 CCT GGG GCT TCA GTG AAG GTT TCC TGC AAG GCT TCT GGA TAC ACA TTC	144														
Pro Gly Ala Ser Val Lys Val Ser Cys Lys Ala Ser Gly Tyr Thr Phe															
15 20 25															
ACT AGT TAT GTT ATT CAC TGG GTG CGA CAG GCC CCT GGT CAG GGC CTT	192														
Thr Ser Tyr Val Ile His Trp Val Arg Gln Ala Pro Gly Gln Gly Leu															
30 35 40 45															

	GAG	TGG	ATG	GGA	TAT	ATT	AAT	CCT	TAC	AAT	GAT	GGG	ACT	AAG	TAC	AAT	240
	Glu	Trp	Met	Gly	Tyr	Ile	Asn	Pro	Tyr	Asn	Asp	Gly	Thr	Lys	Tyr	Asn	
5					50					55					60		
	GAG	AGG	TTC	AAA	GGC	AGA	GTC	ACA	ATC	ACT	GCA	GAC	ACG	TCC	ACG	AGC	288
	Glu	Arg	Phe	Lys	Gly	Arg	Val	Thr	Ile	Thr	Ala	Asp	Thr	Ser	Thr	Ser	
				65					70					75			
10	ACA	GCC	TAC	ATG	GAG	CTC	AGT	TCG	CTG	AGA	TCT	GAG	GAC	ACG	GCG	GTG	336
	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	Leu	Arg	Ser	Glu	Asp	Thr	Ala	Val	
			80					85					90				
15	TAT	CTC	TGT	GGG	AGA	GAA	GGA	ATT	AGG	TAC	TAT	GGT	CTA	CTG	GGA	GAC	384
15	Tyr	Leu	Cys	Gly	Arg	Glu	Gly	Ile	Arg	Tyr	Tyr	Gly	Leu	Leu	Gly	Asp	
		95					100					105					
	TAC	TGG	GGC	CAA	GGC	ACC	CTG	GTC	ACA	GTC	TCC	TCA	G				421
20	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Val	Thr	Val	Ser	Ser					
	110					115					120	121					
	SEQ	ID 1	vo: 6	56													
25	SEQU	JENCE	E LEN	GTH:	96												
	SEQU	JENCE	TYE	PE: r	nucle	eic a	cid										
	STRANDEDNESS: single TOPOLOGY: linear																
00	TOPO	LOGY	(: li	near													
30	MOLE	CULE	TYF	E: C	ther	nuc	cleic	aci	d s	ynth	etic	DNA					
	SEQU	JENCE	DES	CRIE	MOIT	l:											
	AGGC	TTCT	GG A	TACA	CATT	C AC	TAGI	TATO	TTA	TTCA	CTG	GGTG	CGAC	AG G	CCCC	TGGTC	60
35	AGGG	CCTI	GC G	TGGA	TGGG	A TA	TATT	'AATC	CTT	'ACA							96
	SEQ	ID N	10: 6	7													
	SEQU	ENCE	LEN	IGTH:	98												
40	SEQU	ENCE	TYF	e: n	ucle	ic a	cid										
	STRA	NDED	NESS	: si	ngle												
	TOPO	LOGY	: li	near													
	MOLE	CULE	TYP	E: C	ther	nuc	leic	aci	d s	ynth	etic	DNA					
45	SEQU	ENCE	DES	CRIP	TION	:											
	TGTA	.GGCT	GT G	CTCG	TGGA	C CT	GTCT	GCAG	TGA	TTGT	GAC	TCTG	CCTT	TG A	ACCT	CTCAT	60
	TGTA	CTTA	GT C	CCAT	CATT	G TA	AGGA	TTAA	TAT	ATCC	C						96
50																	
	SEQ	ID N	0: 6	8													
	SEOU	ENCE	LEN	GTH:	421												

5

	SEQ	UENC	E TY	PE:	nucl	eic	acid										
	STRANDEDNESS: double																
5	TOP	OLOG	Y: 1	inea	r												
	MOL	ECUL	E TY	PE:	Othe	r nu	clei	c ac	id	synt	heti	C DN	Α				
	MOLECULE TYPE: Other nucleic acid synthetic DNA FEATURE:																
	NAM	E/KE	Y: s	ig p	epti	de											
10	LOC	OITA	N: 1	57													
	IDE	NTIF	ICAT	ION	метн	OD:	S										
	NAM	E/KE	Y: d	omai	n												
15	LOC	ATIO	N: 1	48	162												
10	IDE	NTIF	ICAT:	ION	METH	OD:	s										
	OTH	ER I	NFOR	MATI	ON:	CDR1											
	NAMI	E/KE	Y: d	omai	n												
20	LOC	ATIO	N: 2	05	255												
	IDE	NTIF	ICAT:	ION	METH	OD:	s										
	ОТН	ER I	NFOR	MATI	: NC	CDR2											
	NAM	E/KE	Y: do	omai	n												
25	LOC	OITA	N: 3	52	378												
	IDE	NTIF:	ICAT:	ION I	METH	DD:	S										
	ОТН	ER II	NFOR	MATI	:NC	CDR3											
30	SEQ	JENCI	E DES	SCRI	PTIO	N:											
	ATG	GAA	TGG	AGT	TGG	ATA	TTT	CTC	TTT	CTC	CTG	TCA	GGA	ACT	GCA	GGT	48
	Met	Glu	Trp	Ser	Trp	Ile	Phe	Leu	Phe	Leu	Leu	Ser	Gly	Thr	Ala	Gly	
					-15					-10					-5		
35	GTC	CAC	TCT	GAG	GTC	CAG	CTG	GTG	CAG	TCT	GGA	GCT	GAG	GTG	AAG	AAG	96
	Val	His	Ser	Glu	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys	
			-1	1				5					10				÷
	CCT	GGG	GCT	TCA	GTG	AAG	GTT	TCC	TGC	AAG	GCT	TCT	GGA	TAC	ACA	TTC	144
40	Pro	Gly	Ala	Ser	Val	Lys	Val	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Thr	Phe	
		15					20					25					
	ACT	AGT	TAT	GTT	ATT	CAC	TGG	GTG	CGA	CAG	GCC	CCT	GGT	CAG	GGC	CTT	192
45	Thr	Ser	Tyr	Val	Ile	His	Trp	Val	Arg	Gln	Ala	Pro	Gly	Gln	Gly	Leu	
	30					35					40					45	
	GCG	TGG	ATG	GGA	TAT	ATT	AAT	CCT	TAC	AAT	GAT	GGG	ACT	AAG	TAC	AAT	240
	Ala	Trp	Met	Gly	Tyr	Ile	Asn	Pro	Tyr	Asn	Asp	Gly	Thr	Lys	Tyr	Asn	
50					50					55					60		
	GAG	AGG	TTC	AAA	GGC	AGA	GTC	ACA	ATC	ACT	GCA	GAC	AGG	TCC	ACG	AGC	288
	Glu	Arg	Phe	Lys	Gly	Arg	Val	Thr	Ile	Thr	Ala	Asp	Arg	Ser	Thr	Ser	

	65 70 75	
	ACA GCC TAC ATG GAG CTC AGT TCG CTG AGA TCT GAG GAC ACG GCG GTG	336
5	Thr Ala Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val	
	80 85 90	
	TAT CTC TGT GGG AGA GAA GGA ATT AGG TAC TAT GGT CTA CTG GGA GAC	384
	Tyr Leu Cys Gly Arg Glu Gly Ile Arg Tyr Tyr Gly Leu Leu Gly Asp	
10	95 100 105	
	TAC TGG GGC CAA GGC ACC CTG GTC ACA GTC TCC TCA G	421
	Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Ser	
15	110 115 120 121	
	SEQ ID NO: 69	
	SEQUENCE LENGTH: 96	
20	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
25	SEQUENCE DESCRIPTION:	
	AGGCTTCTGG ATACACATTC ACTAGTTATG TTATTCACTG GGTGCGACAG AGGCCTGGTC	60
	AGGGCCTTGC GTGGATGGGA TATATTAATC CTTACA	96
30		
	SEQ ID NO: 70	
	SEQUENCE LENGTH: 98-7	
	SEQUENCE TYPE: nucleic acid	
35	STRANDEDNESS: single	
	TOPOLOGY: linear	
	MOLECULE TYPE: Other nucleic acid synthetic DNA	
40	SEQUENCE DESCRIPTION:	C 0
40	TGTAGACTGT GCTCGTGGAC CTGTCTGAAG TGATTGTGAC TTTGCCTTTG AACCTCTCAT	60 98
	TGTACTTAGT CCCATCATTG TAAGGATTAA TATATCCC	90
	070 TD NO. 71	
45	SEQ ID NO: 71	
	SEQUENCE LENGTH: 96 SEQUENCE TYPE: nucleic acid	
	~	
	STRANDEDNESS: single TOPOLOGY: linear	
50	MOLECULE TYPE: Other nucleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	

	GTCCACGAGC ACAGTCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA	60													
	TCTCTGTGGG AGAGAAGGAA TTAGGTACTA TGGTCT	96													
5															
	SEQ ID NO: 72														
	SEQUENCE LENGTH: 421														
	SEQUENCE TYPE: nucleic acid														
10	STRANDEDNESS: double														
	TOPOLOGY: linear														
	MOLECULE TYPE: Other nucleic acid synthetic DNA														
	FEATURE:														
15	NAME/KEY: sig peptide														
	LOCATION: 157														
	IDENTIFICATION METHOD: S														
20	NAME/KEY: domain														
	LOCATION: 148162														
	IDENTIFICATION METHOD: S														
	OTHER INFORMATION: CDR1														
25	NAME/KEY: domain														
	LOCATION: 205255														
	IDENTIFICATION METHOD: S														
	OTHER INFORMATION: CDR2														
30	NAME/KEY: domain														
	LOCATION: 352378 -														
	IDENTIFICATION METHOD: S														
35	OTHER INFORMATION: CDR3														
33	SEQUENCE DESCRIPTION:														
	ATG GAA TGG AGT TGG ATA TTT CTC TTT CTC CTG TCA GGA ACT GCA GGT	8													
	Met Glu Trp Ser Trp Ile Phe Leu Phe Leu Leu Ser Gly Thr Ala Gly														
40	· -15 -10 -5														
	GTC CAC TCT GAG GTC CAG CTG GTG CAG TCT GGA GCT GAG GTG AAG AAG 9	6													
	Val His Ser Glu Val Gln Leu Val Gln Ser Gly Ala Glu Val Lys Lys														
	-1 1 5 10														
45	CCT GGG GCT TCA GTG AAG GTT TCC TGC AAG GCT TCT GGA TAC ACA TTC 14	4													
	Pro Gly Ala Ser Val Lys Val Ser Cys Lys Ala Ser Gly Tyr Thr Phe														
	15 20 25														
50	ACT AGT TAT GTT ATT CAC TGG GTG CGA CAG AGG CCT GGT CAG GGC CTT 19	2													
50	Thr Ser Tyr Val Ile His Trp Val Arg Gln Arg Pro Gly Gln Gly Leu														
	30 35 40 45														

	GCG TGG ATG GGA TAT ATT AAT CCT TAC AAT GAT GGG ACT AAG TAC AAT	240												
	Ala Trp Met Gly Tyr Ile Asn Pro Tyr Asn Asp Gly Thr Lys Tyr Asn													
5	50 55 60													
	GAG AGG TTC AAA GGC AAA GTC ACA ATC ACT TCA GAC AGG TCC ACG AGC	288												
	Glu Arg Phe Lys Gly Lys Val Thr Ile Thr Ser Asp Arg Ser Thr Ser													
	65 70 75													
10	ACA GTC TAC ATG GAG CTC AGT TCG CTG AGA TCT GAG GAC ACG GCG GTG	336												
	Thr Val Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val													
	80 85 90													
4.5	TAT CTC TGT GGG AGA GAA GGA ATT AGG TAC TAT GGT CTA CTG GGA GAC	384												
15	Tyr Leu Cys Gly Arg Glu Gly Ile Arg Tyr Tyr Gly Leu Leu Gly Asp													
	95 100 105													
	TAC TGG GGC CAA GGC ACC CTG GTC ACA GTC TCC TCA G	421												
20	Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Ser													
	110 115 120 121													
	SEQ ID NO: 73													
25	SEQUENCE LENGTH: 92													
	SEQUENCE TYPE: nucleic acid													
	STRANDEDNESS: single													
30	TOPOLOGY: linear													
30	MOLECULE TYPE: Other nucleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
	TAGGAGACAG AGTCACCATC ACTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT													
35	GGTATCGGCA GAAACCAGGG AAAGCCCCTG AA	92												
	SEQ ID NO: 74													
	SEQUENCE LENGTH: 90													
40	SEQUENCE TYPE: nucleic acid													
	STRANDEDNESS: single													
	TOPOLOGY: linear													
45	MOLECULE TYPE: other necleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
	TTCCAGACCC GCTGCCACTG AACCTTGATG GGACTCCTGA CTGTAATCTT GATGTGTGT	60												
	AGATCAGGAG TTCAGGGGCT TTCCCTGGTT	90												
50														
	SEQ ID NO: 75													
	SEQUENCE LENGTH: 91													

	SEQUENCE TYPE: nucleic acid														
	STRANDEDNESS: single														
5	TOPOLOGY: linear														
	MOLECULE TYPE: other necleic acid synthetic DNA														
	SEQUENCE DESCRIPTION:														
10	GTTTTCCCAG TCACGACCGT ACGTTTATT TCCACCTTGG TCCCTTGGCC GACCGTGTAC	60													
10	GGAAGCGTAT AACCCTGTTG GCAGTAGTAA G	91													
	SEQ ID NO: 76														
15	SEQUENCE LENGTH: 382														
75	SEQUENCE TYPE: nucleic acid														
	STRANDEDNESS: double														
	TOPOLOGY: linear														
20	MOLECULE TYPE: other necleic acid synthetic DNA FEATURE:														
	NAME/KEY: sig peptide														
	LOCATION: 160														
25	IDENTIFICATION METHOD: S														
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	IDENTIFICATION METHOD: S														
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	LOCATION: 325351														
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45	Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln														
	-20 -15 -10 -5														
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50	Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser														
	-1 1 5														
	GCC TCT GTA GGA GAC AGA GTC ACC ATC ACT TGC GGG ACA AGT GAG GAC 14	44													

	Ala Ser Val Gly Asp Arg Val Thr Ile Thr Cys Gly Thr Ser Glu Asp														
	15 20 25														
5	ATT ATC AAT TAT TTA AAC TGG TAT CGG CAG AAA CCA GGG AAA GCC CCT	192													
	Ile Ile Asn Tyr Leu Asn Trp Tyr Arg Gln Lys Pro Gly Lys Ala Pro														
	30 35 40														
10	GAA CTC CTG ATC TAC CAC ACA TCA AGA TTA CAG TCA GGA GTC CCA TCA	240													
10	Glu Leu Leu Ile Tyr His Thr Ser Arg Leu Gln Ser Gly Val Pro Ser														
	45 50 55 60														
	AGG TTC AGT GGC AGC GGG TCT GGA ACA GAT TTC ACT CTC ACC ATT AGT	288													
15	Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser														
	65 70 75														
	AGT CTG CAA CCT GAA GAT TTT GCC ACT TAC TAC TGC CAA CAG GGT TAT	336													
	Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Gly Tyr														
20	80 85 90														
	ACG CTT CCG TAC ACG GTC GGC CAA GGG ACC AAG GTG GAA ATA AAA C	382													
	Thr Leu Pro Tyr Thr Val Gly Gln Gly Thr Lys Val Glu Ile Lys														
25	95 100 105 107														
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	SEQUENCE LENGTH: 92														
30	SEQUENCE TYPE: nucleic acid														
	STRANDEDNESS: single														
	TOPOLOGY: linear														
35	MOLECULE TYPE: other necleic acid synthetic DNA														
	SEQUENCE DESCRIPTION:														
	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT														
	GGTATCGGCA GAAACCAGGG AAAGCCCCTG AA	92													
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	SEQ ID NO: 78														
	SEQUENCE LENGTH: 88														
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	STRANDEDNESS: single														
	TOPOLOGY: linear														
	MOLECULE TYPE: other necleic acid synthetic DNA														
50	SEQUENCE DESCRIPTION:														
	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTGACCTGC AACCTGAAGA	60													
	TTTTGCCACT TACTACTGCC AACAGGGT	88													

	SEC) ID	NO:	79													
	SEC	QUENC	CE LI	ENGT	H: 38	32											
5	SEC	QUENC	E T	PE:	nuc	leic	acio	i									
	STF	RANDE	EDNES	SS: d	doub	le											
	TOF	OLOG	Y: 1	line	ar												
	MOI	ECUI	E T	PE:	othe	er ne	eclei	ic ac	id	synt	heti	LC D	IA.				
10	FEA	TURE	Ξ:														
	NAM	IE/KE	Y: 5	sig p	pepti	lde											
	LOC	CATIC	N: 1	60)												
15	IDE	NTIF	CAT	NOI	METH	OD:	S										
10	NAM	E/KE	Y: d	lomai	.n												
	LOC	ATIO	N: 1	.30	162												
	IDENTIFICATION METHOD: S																
20	OTHER INFORMATION: CDR1																
	NAME/KEY: domain																
	LOC	ATIO	N: 2	08	228												
25	IDE	NTIF	ICAT	ION	METH	OD:	S										
	OTH	ER I	NFOR	MATI	ON:	CDR2											
	NAM	E/KE	Y: d	omai	n												
	LOCATION: 325351																
30	IDENTIFICATION METHOD: S																
30	OTHER INFORMATION: CDR3 SEQUENCE DESCRIPTION:																
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	ATG	ATG	TCC	TCT	GCT	CAG	TTC	CTT	GGT	CTC	CTG	TTG	CTC	TGT	TTT	CAA	48
35	Met	Met	Ser	Ser	Ala	Gln	Phe	Leu	Gly	Leu	Leu	Leu	Leu	Cys	Phe	Gln	
	-20					-15					-10					-5	
	GAT	ATC	AGA	TGT	GAT	ATC	CAG	ATG	ACA	CAG	TCT	CCA	TCC	TCC	CTG	TCT	. 96
	Asp	Ile	Arg	Cys	Asp	Ile	Gln	Met	Thr	Gln	Ser	Pro	Ser	Ser	Leu	Ser	
40				·-1	1				5					10			
	GCC	TCT	GTA	GGA	GAC	AGA	GTC	ACC	ATC	GGT	TGC	GGG	ACA	AGT	GAG	GAC	144
	Ala	Ser	Val	Gly	Asp	Arg	Val	Thr	Ile	Gly	Cys	Gly	Thr	Ser	Glu	Asp	
45			15					20					25				
45	ATT	ATC	AAT	TAT	TTA	AAC	TGG	TAT	CGG	CAG	AAA	CCA	GGG	AAA	GCC	CCT	192
	Ile	Ile	Asn	Tyr	Leu	Asn	Trp	Tyr	Arg	Gln	Lys	Pro	Gly	Lys	Ala	Pro	
		30					35					40					
50	GAA	CTC	CTG	ATC	TAC	CAC	ACA	TCA	AGA	TTA	CAG	TCA	GGA	GTC	CCA	TCA	240
	Glu	Leu	Leu	Ile	Tyr	His	Thr	Ser	Arg	Leu	Gln	Ser	Gly	Val	Pro	Ser	
	45					50					55					60	

	AGG TTC AGT GGC AGC GGG TCT GGA ACA GAT TTC ACT CTC ACC ATT AGT	288												
	Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser													
5	65 70 75													
	GAC CTG CAA CCT GAA GAT TTT GCC ACT TAC TAC TGC CAA CAG GGT TAT	336												
	Asp Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Gly Tyr													
	80 85 90													
10	ACG CTT CCG TAC ACG GTC GGC CAA GGG ACC AAG GTG GAA ATA AAA C	382												
	Thr Leu Pro Tyr Thr Val Gly Gln Gly Thr Lys Val Glu Ile Lys													
	95 100 105 107													
45														
15	SEQ ID NO: 80													
	SEQUENCE LENGTH: 83													
	SEQUENCE TYPE: nucleic acid													
20	STRANDEDNESS: single													
	TOPOLOGY: linear													
	MOLECULE TYPE: other necleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
25	GATGGTGACT CTGTCTCCTA CAGAGGCAGA CAGGGAGGAT GTAGCCTGTG TCATCTGGAT	60												
	ATCACATCTG ATGTCTTGAA AAC	23												
30	SEQ ID NO: 81													
30	SEQUENCE LENGTH: 92													
	SEQUENCE TYPE: nucleïc acid													
	STRANDEDNESS: single													
35	TOPOLOGY: linear													
	MOLECULE TYPE: other necleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60												
40	GGTATCGGAA GAAACCAGGG AAAGCCCCTG AA	92												
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45	SEQUENCE LENGTH: 88													
	SEQUENCE TYPE: nucleic acid													
	STRANDEDNESS: single													
	TOPOLOGY: linear													
50	MOLECULE TYPE: other necleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTGACCTGC AACCTGAAGA	60												

	TTTTGCCACT TACTTTTGCC AACAGGGT	88												
5	SEQ ID NO: 83													
	SEQUENCE LENGTH: 91													
	SEQUENCE TYPE: nucleic acid													
	STRANDEDNESS: single													
10	TOPOLOGY: linear													
	MOLECULE TYPE: other necleic acid synthetic DNA													
	SEQUENCE DESCRIPTION:													
15	GTTTTCCCAG TCACGACCGT ACGTTTTATT TCCACCTTGG TCCCTTGGCC GACCGTGTAC	60												
	GGAAGCGTAT AACCCTGTTG GCAAAAGTAA G	91												
	SEQ ID NO: 84													
20	SEQUENCE LENGTH: 382													
	SEQUENCE TYPE: nucleic acid													
	STRANDEDNESS: double													
	TOPOLOGY: linear													
25	MOLECULE TYPE: other necleic acid synthetic DNA													
	FEATURE:													
	NAME/KEY: sig peptide													
	LOCATION: 160													
30	IDENTIFICATION METHOD: S													
	NAME/KEY: domain													
	LOCATION: 130162													
35	IDENTIFICATION METHOD: S													
	OTHER INFORMATION: CDR1													
	NAME/KEY: domain													
	LOCATION: 208228													
40	IDENTIFICATION METHOD: S													
	OTHER INFORMATION: CDR2													
	NAME/KEY: domain													
45	LOCATION: 325351													
45	IDENTIFICATION METHOD: S													
	OTHER INFORMATION: CDR3													
	SEQUENCE DESCRIPTION:													
50	ATG ATG TCC TCT GCT CAG TTC CTT GGT CTC CTG TTG CTC TGT TTT CAA	48												
	Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln													
	-20 -15 -10 -5													

	GAT ATC AGA TGT GAT ATC CAG ATG ACA CAG GCT ACA TCC TCC CTG TCT 96
	Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ala Thr Ser Ser Leu Ser
5	-1 1 5 10
	GCC TCT GTA GGA GAC AGA GTC ACC ATC GGT TGC GGG ACA AGT GAG GAC 144
	Ala Ser Val Gly Asp Arg Val Thr Ile Gly Cys Gly Thr Ser Glu Asp
10	15 20 25
70	ATT ATC AAT TAT TTA AAC TGG TAT CGG AAG AAA CCA GGG AAA GCC CCT 192
	Ile Ile Asn Tyr Leu Asn Trp Tyr Arg Lys Lys Pro Gly Lys Ala Pro
	30 35 40
15	GAA CTC CTG ATC TAC CAC ACA TCA AGA TTA CAG TCA GGA GTC CCA TCA 240
	Glu Leu Leu Ile Tyr His Thr Ser Arg Leu Gln Ser Gly Val Pro Ser 45 50 55 60
	45 50 55 60 AGG TTC AGT GGC AGC GGG TCT GGA ACA GAT TTC ACT CTC ACC ATT AGT 288
00	Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser
20	65 70 75
	GAC CTG CAA CCT GAA GAT TTT GCC ACT TAC TTT TGC CAA CAG GGT TAT 336
	Asp Leu Gln Pro Glu Asp Phe Ala Thr Tyr Phe Cys Gln Gln Gly Tyr
25	80 85 90
	ACG CTT CCG TAC ACG GTC GGC CAA GGG ACC AAG GTG GAA ATA AAA C 382
	Thr Leu Pro Tyr Thr Val Gly Gln Gly Thr Lys Val Glu Ile Lys
20	95 100 105
30	
	SEQ ID NO: 85
	SEQUENCE LENGTH: 92
35	SEQUENCE TYPE: nucleic acid
	STRANDEDNESS: single
	TOPOLOGY: linear
40	MOLECULE TYPE: other necleic acid synthetic DNA
40	SEQUENCE DESCRIPTION:
	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT 60
	GGTATCGGAA GAAACCAGGG AAAGCCGTTG AA 92
45	SEQ ID NO: 86
	SEQUENCE LENGTH: 90
	SEQUENCE TYPE: nucleic acid
50	STRANDEDNESS: single
50	TOPOLOGY: linear
	MOLECULE TYPE: other necleic acid synthetic DNA
	•

	SEQUENCE DESCRIPTION:	
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5	AGATCAGGAG TTCAACGGCT TTCCCTGGTT	90
	SEQ ID NO: 87	
	SEQUENCE LENGTH: 88	
10	SEQUENCE TYPE: nucleic acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
15	MOLECULE TYPE: other necleic acid synthetic DNA	
15	SEQUENCE DESCRIPTION:	
	CAGTGGCAGC GGGTCTGGAA CAGATTATAC TCTCACCATT AGTGACCTGC AACCTGAAGA	60
	TTTTGCCACT TACTTTTGCC AACAGGGT	88
20	SEO ID NO: 88	
	SEQUENCE LENGTH: 382	
	SEQUENCE TYPE: nucleic acid	
25	STRANDEDNESS: double	
20	TOPOLOGY: linear	
	MOLECULE TYPE: other necleic acid synthetic DNA	
	FEATURE:	
30	NAME/KEY: sig peptide	
	LOCATION: 160	
	IDENTIFICATION METHOD: S	
	NAME/KEY: domain	
35	LOCATION: 130162	
	IDENTIFICATION METHOD: S	
	OTHER INFORMATION: CDR1	
40	NAME/KEY: domain	
	LOCATION: 208228	
	IDENTIFICATION METHOD: S	
	OTHER INFORMATION: CDR2	
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	LOCATION: 325351	
	IDENTIFICATION METHOD: S	
50	OTHER INFORMATION: CDR3	
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5	GAT	ATC	AGA	TGT	GAT	ATC	CAG	ATG	ACA	CAG	GCT	ACA	TCC	TCC	CTG	TCT	96
	Asp	Ile	Arg	Cys	Asp	Ile	Gln	Met	Thr	Gln	Ala	Thr	Ser	Ser	Leu	Ser	
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10	GCC	TCT	GTA	GGA	GAC	AGA	GTC	ACC	ATC	GGT	TGC	GGG	ACA	AGT	GAG	GAC	144
10	Ala	Ser	Val	Gly	Asp	Arg	Val	Thr	Ile	Gly	Cys	Gly	Thr	Ser	Glu	Asp	
			15					20					25				
					TTA												192
15	Ile		Asn	Tyr	Leu	Asn	_	Tyr	Arg	Lys	Lys		Gly	Lys	Ala	Val	
		30					35					40					
					TAC												240
		Leu	Leu	IIe	Tyr		Tnr	Ser	Arg	Leu		Ser	GIĀ	vai	Pro		
20	45	ጥጥር	አ ርጥ	ccc	AGC	50	m/m	CCA	A C A	<i>C</i> አጥ	55 Tat	አ ርጥ	CTC	N.C.C	א חייחי	60	288
					Ser												200
	ur 9	FILE	Ser	Gry	65	GIY	561	GIY	1111	70	111	1111	пеа	1111	75	Ser	
25	GAC	CTG	CAA	ССТ	GAA	GAT	ттт	GCC	ACT		ттт	TGC	CAA	CAG		TAT	336
					Glu												
	-			80		•			85	•		•		90	•	-	
	ACG	CTT	CCG	TAC	ACG	GTC	GGC	ÇAA	GGG	ACC	AAG	GTG	GAA	ATA	AAA	С	382
30	Thr	Leu	Pro	Tyr	Thr	Val	Gly	Gln	Gly	Thr	Lys	Val	Gľu	Ile	Lys		
			95					100					105		107		
35	SEQ	ID N	io: 8	19													
	SEQU	ENCE	LEN	IGTH:	25												
	SEQU	ENCE	TYP	E: n	ucle	ic a	cid										
	STRA	NDED	NESS	: si	.ngle	:											
40	TOPO	LOGY	: li	near	•												
	MOLE	CULE	TYP	E: 0	ther	nec	leic	aci	d s	ynth	etic	DNA	,				
	SEQU	ENCE	DES	CRIP	TION	·:											
45	GCTT	CCAC	CA A	.GGGC	CCAT	C CG	TCT										25
	SEQ																
	SEQU																
50	SEQU						Cld										
	STRA				-												
	TOPOLOGY: linear																

	MOLECULE TYPE: other necleic acid synthetic DNA	
	SEQUENCE DESCRIPTION:	
5	AAGGATCCTG GCACTCATTT ACCCAGAGAC	30
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40	SEQUENCE LENGTH: 313	
10	SEQUENCE TYPE: amino acid	
	STRANDEDNESS: single	
	TOPOLOGY: linear	
15	MOLECULE TYPE: protein	
	SEQUENCE DESCRIPTION:	
	Asp Leu Leu Pro Asp Glu Lys Ile Ser Leu Leu Pro Pro Val Asn	
20	1 5 10 15	
20	Phe Thr Ile Lys Val Thr Gly Leu Ala Gln Val Leu Leu Gln Trp	
	20 25 30	
25	Lys Pro Asn Pro Asp Gln Glu Gln Arg Asn Val Asn Leu Glu Tyr	
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	Gln Val Lys Ile Asn Ala Pro Lys Glu Asp Asp Tyr Glu Thr Arg	
30	50 55 60	
	Ile Thr Glu Ser Lys Çys Val Thr Ile Leu His Lys Gly Phe Ser	
	65 70 75	
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35	Ala Ser Val Arg Thr Ile Leu Gln Asn Asp His Ser Leu Leu Ala	
	80 85 90	
	Ser Ser Trp Ala Ser Ala Glu Leu His Ala Pro Pro Gly Ser Pro	
40	. 95 100 105	
	Gly Thr Ser Val Val Asn Leu Thr Cys Thr Thr Asn Thr Thr Glu	
	110 115 120	
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	Asp Asn Tyr Ser Arg Leu Arg Ser Tyr Gln Val Ser Leu His Cys	
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	Leu Tyr Tyr Arg Tyr Gly Ser Trp Thr Glu Glu Cys Gln Glu Tyr	

			155			160		165
5	Ser Lys	Asp Thr	Leu Gl	y Arg A	Asn Ile	Ala Cys	Trp Phe	Pro Arg
5			170			175		180
	Thr Phe	Ile Leu	Ser Ly	s Gly A	Arg Asp	Trp Leu	Ala Val	Leu Val
10			185			190		195
	Asn Gly	Ser Ser	Lys Hi	s Ser A	Ala Ile	Arg Pro	Phe Asp	Gln Leu
			200			205		210
15	Phe Ala	Leu His	Ala Il	e Asp G	In Ile	Asn Pro	Pro Leu	Asn Val
			215		:	220		225
20	Thr Ala	Glu Ile	Glu Gl	y Thr A	rg Leu	Ser Ile	Gln Trp	Glu Lys
			230		:	235		240
	Pro Val	Ser Ala	Phe Pro	o Ile H	lis Cys I	Phe Asp	Tyr Glu	Val Lys
25			245		;	250		255
	Ile His	Asn Thr	Arg Ası	n Gly T	yr Leu (Gln Ile	Glu Lys	Leu Met
			260		7	265		270
30	Thr Asn	Ala Phe	Ile Se	r Ile I	le Asp A	Asp Leu	Ser Lys	Tyr Asp
			275		2	280		285
35	Val Gln	Val Arg	Ala Ala	a Val S	er Ser M	Met Cys	Arg Glu	Ala Gly
			290		2	295		300
	Leu Trp	Ser Glu	Trp Ser	Gln P	ro Ile T	Cyr Val	Gly Lys	
40			305		3	310	313	

SEQUENCE LISTING

5	(1) GENERAL INFORMATION:	
10	(i) APPLICANT: (A) NAME: Kyowa Hakko Kogyo Co., Ltd. (B) STREET: 6-1, Ohtemachi 1-chome (C) CITY: Chiyoda-ku (D) STATE: Tokyo (E) COUNTRY: Japan (F) POSTAL CODE (ZIP): 100	
15	(ii) TITLE OF INVENTION: Antibody against human interleukin-5 rece α chain	ptoi
	(iii) NUMBER OF SEQUENCES: 106	
20	 (iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO) 	
25	(2) INFORMATION FOR SEQ ID NO: 1:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
30	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
	CAAAGCTTAC CATGATCATC GTGGCGCATG TA	32
40	(2) INFORMATION FOR SEQ ID NO: 2:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 32 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single	
45	(D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
50	(iii) HYPOTHETICAL: YES	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:	
	CAGGATCCCT ACTTACCCAC ATAAATAGGT TG	32

90

	(2) INFORMATION FOR SEQ ID NO: 3:	
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:	
15	CAGATATCTC ACTTCTCCCA CCTGTCA	2.
	(2) INFORMATION FOR SEQ ID NO: 4:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
	AGCTTCCACC ATGGAGTTTG GGCTCAGCTG GCTTTTTCTT GTCCTTGTTT TCAAAGGTGT	60
<i>35</i>	TCAGTGTGAC TTACTTCCTG ATGAAAAG	88
	(2) INFORMATION FOR SEQ ID NO: 5:	
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 84 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
50	CTTTTCATCA GGAAGTAAGT CACACTGAAC ACCTTTGAAA ACAAGGACAA GAAAAAGCCA	60
	GCTGAGCCCA AACTCCATGG TGGA	84

	(2) INFORMATION FOR SEQ ID NO: 6:	
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 51 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
	AGCTTCCACC ATGGCTACAG GCTCCCGGAC GTCCCTGCTC CTGGCTTTTG G	51
20	(2) INFORMATION FOR SEQ ID NO: 7:	
20	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 58 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
30	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
	CCTGCTCTGC CTGCCCTGGC TTCAAGAGGG CAGTGCCGAC TTACTTCCTG ATGAAAAG	58
35	(2) INFORMATION FOR SEQ ID NO: 8:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 64 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
45	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:	
50	CTTTTCATCA GGAAGTAAGT CGGCACTGCC CTCTTGAAGC CAGGGCAGGC AGAGCAGGCC	60
	AAAA	64

	(2) INFORMATION FOR SEQ ID NO: 9:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:	
15	GCCAGGAGCA GGGACGTCCG GGAGCCTGTA GCCATGGTGG A	41
	(2) INFORMATION FOR SEQ ID NO: 10:	
20	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
25	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
30	GGCAGCGGCG GTTCCGGTGA GCCCAAATCT TGTGACAAA	39
	(2) INFORMATION FOR SEQ ID NO: 11:	
35	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 34 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
40	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
45	CAAAGCTTCC ACCATGGAGT TTGGGCTCAG CTGG	34
	(2) INFORMATION FOR SEQ ID NO: 12:	
50	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 34 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single	

	(D) TOPOLOGY: linear	
5	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
10	CAAAGCTTCC ACCATGGAGT TTGGGCTCAG CTGG	34
	(2) INFORMATION FOR SEQ ID NO: 13:	
15	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
20	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	
25	ACCGGAACCG CCGCTGCCCT TACCCACATA AATAGGTTG	39
	(2) INFORMATION FOR SEQ ID NO: 14:	
30	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 34 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
35	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
40	CAAAGCTTCC ACCATGGCTA CAGGCTCCCG GACG	34
	(2) INFORMATION FOR SEQ ID NO: 15:	
45	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 76 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: double(D) TOPOLOGY: linear	
50	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
	CGATAAGCTA TGAAAACTAC AGCCTTGGAG GAAGCTTAAA TGAGCTCGAT ATCAAGGCCT	60
5	ACCCGGGCGC CATGCA	76
	(2) INFORMATION FOR SEQ ID NO: 16:	
10	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
15	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	
	CACTCAGTGT TAACTGAGGA GCAGGTGAAT TC	32
	(2) INFORMATION FOR SEQ ID NO: 17:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 40 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
30	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:	
	AGCTGAATTC ACCTGCTCCT CAGTTAACAC TGAGTGGTAC	40
40	(2) INFORMATION FOR SEQ ID NO: 18:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
45	(ii) MOLECULE TYPE: other nucleic acid	
	(A) DESCRIPTION: /desc = "synthetic DNA"	
50	(iii) HYPOTHETICAL: YES	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:	
	AATTCGTACG GTGGCTGCAC C	21

	(2) INFORMATION	FOR SEQ ID NO: 19:	
5	(A) LE (B) TY (C) ST	CE CHARACTERISTICS: ENGTH: 17 base pairs YPE: nucleic acid FRANDEDNESS: single DPOLOGY: linear	
10		LE TYPE: other nucleic acid ESCRIPTION: /desc = "synthetic DNA"	
	(iii) HYPOTHE	ETICAL: YES	
15	(xi) SEQUENCE	CE DESCRIPTION: SEQ ID NO: 19:	
	GGTGCAGCCA CCGTA	ACG	17
20	(2) INFORMATION 1	FOR SEQ ID NO: 20:	
25	(A) LEI (B) TYI (C) STI	CE CHARACTERISTICS: CNGTH: 26 base pairs YPE: nucleic acid CRANDEDNESS: single OPOLOGY: linear	
		E TYPE: other nucleic acid SCRIPTION: /desc = "synthetic DNA"	
30	(iii) HYPOTHE	TICAL: YES	
	(xi) SEQUENCE	E DESCRIPTION: SEQ ID NO: 20:	
05	CTCGCGACTA GTGGGC	CCCGC GGCCGC	26
35	(2) INFORMATION E	FOR SEQ ID NO: 21:	
40	(A) LEN (B) TYP (C) STR	E CHARACTERISTICS: NGTH: 34 base pairs PE: nucleic acid RANDEDNESS: single POLOGY: linear	
45	*	E TYPE: other nucleic acid SCRIPTION: /desc = "synthetic DNA"	
	(iii) HYPOTHET	TICAL: YES	
50	(xi) SEQUENCE	E DESCRIPTION: SEQ ID NO: 21:	
50	AGCTGCGGCC GCGGGC	CCCAC TAGTCGCGAG GTAC	34

	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:	22:								
5		(i	(A) L B) T C) S	ENGT YPE : TRAN	HARA H: 4 nuc DEDN OGY:	21 b leic ESS:	ase aci dou	pair d	s							
10		(ii) MO	LECU	LE T	YPE:	CDN	Α									
		(iii) НҮ	POTH	ETIC	AL:	YES										
		(ix		ATUR A) N		KEY:	CDS										
15						ION:	14	20									
		(ix	(.		AME/	KEY:			tide								
20																	
		(xi) SE	QUEN	CE D	ESCR	IPTI(ON:	SEQ	ID N	0: 2	2:					
														TTA Leu		GGT Gly	48
25	1				5					10					15	•	
														TTA Leu		AAG	96
	vai	O.I.I	Cyb	20	•	O.I.I.	J.Cu	Vu	25	501	Gly	GLY	veb	30	val	пув	
30														TTC			144
	PIO	GIŞ	35	ser	ьeu	гуя	Leu	ser 40	Cys	AIA	Ala	ser	45	Phe	Thr	Phe	
				_		_								AAG			192
35	261	50	IYL	GIY	Mec	ALG	55	116	Arg	GIII	116	60	ASP	Lys	AIG	PIO	
														CAC			240
	GIu 65	Trp	Val	Ala	Ala	70	Ser	Ser	GIY	GIŸ	Ser 75	Tyr	Ile	His	Phe	Pro 80	
40														GCC			288
	Asp	ser	Leu	гуѕ	85 85	Arg	Pne	Tnr	vai	90	Arg	Asp	Asn	Ala	ьув 95	Asn	
														ACA Thr			336
45	1111	Leu	ıyı	100	Giu	Mec	261	GIY	105	цуъ	261	GIU	Asp	110	Ala	Met	
														GCT			384
	ıyr	ıyr	115	AIA	Arg	wrg	GTÅ	120	TÀL	GТĀ	ASII	īĀĻ	125	Ala	мес	Asp	
50		TGG											G				421
	TYT	Trp 130	стА	GIII	GTÀ	THE	135	vaı	Inf	val	ser.	140					

	(2)	INF	ORMA	TION	FOR	SEQ	ID :	NO:	23:							
5			()	A) L B) T	ENGT YPE :	CHA H: 1 ami OGY:	40 a	mino cid								
10						YPE: ESCR			SEQ :	ID N	0: 2	3:				
, •	Met 1	Asn	Phe	Gly	Leu 5	Ser	Leu	Ile	Phe	Leu 10	Ala	Leu	Ile	Leu	Lys 15	Gly
15	Val	Gln	Cys	Glu 20	Val	Gln	Leu	Val	Glu 25	Ser	Gly	Gly	Asp	Leu 30	Val	Lys
	Pro	Gly	Gly 35	Ser	Leu	Lys	Leu	Ser 40	Суз	Ala	Ala	Ser	Gly 45	Phe	Thr	Phe
20	Ser	As p 50	Tyr	Gly	Met	Ala	Trp 55	Ile	Arg	Gln	Ile	Ser 60	Asp	Lys	Arg	Pro
	Glu 65	Trp	Val	Ala	Ala	Ile 70	Ser	Ser	Gly	Gly	Ser 75	Tyr	Ile	His	Phe	Pro 80
25	Asp	Ser	Leu	Lys	Gly 85	Arg	Phe	Thr	Val	Ser 90	Arg	Asp	Asn	Ala	Lys 95	Asn
	Thr	Leu	Tyr	Leu 100	Glu	Met	Ser	Gly	Leu 105	Lys	Ser	Glu	Asp	Thr 110	Ala	Met
30	Tyr	Tyr	Cys 115	Ala	Arg	Arg	Gly	Phe 120	Tyr	Gly	Asn	Tyr	Arg 125	Ala	Met	Asp
	Tyr	Trp 130	Gly	Gln	Gly	Thr	Ser 135	Val	Thr	Val	Ser	Ser 140				
35	(2)	TNFC	гамас	יא	FOR	SEQ	א מד	IO · 2	ο4 -							
40	(2)		SEQ (F (E	UENC A) LE B) TY	CE CH ENGTH PE: TRANI	IARAC I: 39 nucl EDNE	TERI 04 ba eic ESS:	STIC use p acid	S: airs	ı						
		(ii)	MOI	ECUI	E TY	PE:	CDNA									
45	((iii)	HYE	POTHE	TICA	L: Y	ES									
50		(ix)		A) NA	ME/K	EY:		13								
		(ix)) NA	ME/K	EY: ON:1			ide							

		(xi) SE	QUEN	CE D	ESCR:	IPTI	ON:	SEQ :	ID N	0: 2	4:					
5	ATG Met	GAG Glu	AAA Lys	GAC Asp	ACA Thr 145	CTC Leu	CTG Leu	CTA Leu	TGG Trp	GTC Val 150	CTG Leu	CTT Leu	CTC Leu	TGG Trp	GTT Val 155	CCA Pro	4.8
				AGT Ser 160													96
10				GGG Gly													144
15				AAT Asn													192
20				CCC Pro													240
20				GCC Ala													288
25				CAT His 240													336
30				AAG Lys													384
		ATC Ile 270		С													394
35	(2)			TION SEQUE													
40			(<i>I</i>	A) LE B) TY D) TO	NGTH PE:	I: 13 amin	l am	nino cid									
				LECUI			_		EQ I	D NC): 25	5:					
45	Met 1	Glu	Lys	Asp	Thr 5	Leu	Leu	Leu	Trp	Val 10	Leu	Leu	Leu	Trp	Val 15	Pro	
	Gly	Ser	Arg	Ser 20	Asp	Ile	Val	Leu	Thr 25	Gln	Ser	Pro	Ala	Ser 30	Leu	Ala	
50	Val	Ser	Leu 35	Gly	Gln	Arg	Ala	Thr 40	Ile	Ser	Cys	Arg	Ala 45	Asn	Glu	Ser	
	Val	Asp 50	His	Asn	Gly	Val	Asn 55	Phe	Met	Asn	Trp	Phe 60	Gln	Gln	Lys	Pro	

	Gly 65		Ser	Pro	Lys	Leu 70	Leu	Ile	Tyr	Ala	Ala 75	Ser	Asn	Gln	Gly	Ser 80	
5	Gly	Val	Pro	Ala	Arg 85	Phe	Ser	Gly	Ser	Gly 90		Gly	Thr	Asp	Phe 95	Ser	
	Leu	Asn	Ile	His 100	Pro	Met	Glu	Glu	Asp 105	Asp	Ala	Ala	Met	Tyr 110	Phe	Сув	
10	Gln	Gln	Ser 115	Lys	Asp	Val	Pro	Trp 120		Phe	Gly	Gly	Gly 125	Thr	Arg	Leu	
	Glu	Ile 130	Lys														
15	(2)	INF	ORMA'	rion	FOR	SEQ	ID 1	NO: :	26:								
20		(i)	() ()	QUENCA) LE 3) T'S C) S'I	engti (PE : [Rani	H: 42 nucl	21 ba leic ESS:	ase p acio doul	pair: d	5							
		(ii)	MOI	ECUI	E T	PE:	CDN	A									
25		(iii)	HYI	POTHE	ETIC	AL: Y	ÆS										
		(ix)		ATURE A) NA B) LC	ME/F			20									
30		(ix)		ATURE A) NA B) LC	ME/F				ide								
35		(xi)	SEÇ	UENC	E DE	ESCRI	PTIC	on: s	SEQ I	D NO): 26	5:					
			TGG Trp														48
40			TCT Ser 150														96
45			GCT Ala														144
			TAT Tyr														192
50			ATT Ile														240

	GAG Glu	AGG Arg	TTC Phe	AAA Lys 215	GGC Gly	AAG Lys	GCC Ala	ACA Thr	CTG Leu 220	ACT Thr	TCA Ser	GAC Asp	AGA Arg	TCC Ser 225	TCC Ser	AGC Ser	288
5				Met	GAG Glu											GTC Val	336
10	TAT Tyr	CTC Leu 245	TGT Cys	GGG Gly	AGA Arg	GAA Glu	GGA Gly 250	ATT Ile	AGG Arg	TAC Tyr	TAT Tyr	GGT Gly 255	CTA Leu	CTG Leu	GGA Gly	GAC Asp	384
15					GGC Gly								G				421
	(2)	INFO	ORMA'	rion	FOR	SEQ	ID 1	NO: :	27:								
20		·	(1	A) Li 3) T	ENCE ENGTI YPE : OPOLO	H: 14 amir	10 ar	mino cid									
25					LE TY		_		SEQ 1	ID NO	D: 27	' :					
	Met 1	Glu	Trp	Ser	Trp 5	Ile	Phe	Leu	Phe	Leu 10	Leu	Ser	Gly	Thr	Ala 15	Gly	
30	Val	His	Ser	Glu 20	Val	Gln	Leu	Gln	Gln 25	Ser	Gly	Pro	Glu	Leu 30	Val	Lys	
35	Pro	Gly	Ala 35	Ser	Val	Lys	Met	Ser 40	Cys	Lys	Ala	Ser	Gly 45	Tyr	Thr	Phe	
	Thr	Ser 50	Tyr	Val	Ile	His	Trp 55	Val	Lys	Gln	Arg	Pro 60	Gly	Gln	Gly	Leu	
40	Ala 65	Trp	Ile	Gly	Tyr	Ile 70	Asn	Pro	Tyr	Asn	Asp 75	Gly	Thr	Lys	Tyr	Asn 80	
	Glu	Arg	Phe	Lys	Gly 85	Lys	Ala	Thr	Leu	Thr 90	Ser	Asp	Arg	Ser	Ser 95	Ser	
45	Thr	Val	Tyr	Met 100	Glu	Leu	Ser	Ser	Leu 105	Thr	Ser	Glu	Asp	Ser 110	Ala	Val	
	Tyr	Leu	Cys 115	Gly	Arg	Glu	Gly	Ile 120	Arg	Tyr	Tyr	Gly	Leu 125	Leu	Gly	Asp	
50	По	Trn	Clar	C1 5	C1	Thr	Thr	T.e.11	Thr	Va l	Sar	ear					

	(2)	INF	orma	TION	FOR	SEQ	ID	NO:	28:								
5		(i	(. (:	A) L B) T C) S	CE C ENGT YPE: TRAN	H: 3 nuc DEDN	82 b leic ESS:	ase aci dou	pair d	ន							
10		(ii)) MO	LECU	LE T	YPE:	cDN.	A									
10		(iii)) НҮ:	POTH:	ETIC	AL:	YES										
15		(ix)	()		E: AME/I OCAT:			81									
20		(ix)	(2		E: AME/1 OCAT:				tide								
		(xi)	SE	QUEN	CE DI	ESCR:	IPTI	эи: :	SEQ :	ID N	D: 2	В:					
		ATG															48
25	Met	Met	Ser	Ser	Ala 145	Gln	Phe	Leu	Gly	Leu 150	Leu	Leu	Leu	Cys	Phe 155	Gln	
		ATC															96
	Asp	Ile	Arg	160	Asp	ire	GIII	мес	165	GIII	Ala	Thr	ser	170	Leu	ser	
30	GCC	TCT	ርፐር	GGA	GAC	AGA	GTC	ልሮሮ	ልጥሮ	GGT	ፐርር	GGG	מרמ	ልርጥ	GAG	GAC	144
		Ser															111
35		ATC															192
	Ile	Ile 190	Asn	Tyr	Leu	Asn	Trp 195	Tyr	Arg	Lys	Lys	Pro 200	Asp	Gly	Thr	Val	
		CTC															240
40	205	Leu				210					215		-			220	
		TTC Phe															288
	3				225	•				230	•				235		
45		CTG															336
	Asp	Leu	Glu	Gln 240	Glu	Asp	Ile	Ala	Thr 245	Tyr	Phe	Cys	Gln	Gln 250	Gly	Tyr	
50		CTT Leu															381
50			255	- 4 -			- - 4	260	- 4				265				
	C																382

	(2)	INF	ORMA	TION	FOR	SEQ	ID :	NO:	29:							
5			(1	A) L B) T	ENCE ENGT YPE : OPOL	H: 1: ami:	27 ai	mino cid								
10) MO							ID N	0: 2:	9:				
	Met 1	Met	Ser	Ser	Ala 5	Gln	Phe	Leu	Gly	Leu 10	Leu	Leu	Leu	Cys	Phe 15	Gln
15	Asp	Ile	Arg	Cys 20	Asp	Ile	Gln	Met	Thr 25	Gln	Ala	Thr	Ser	Ser 30	Leu	Ser
	Ala	Ser	Leu 35	Gly	Asp	Arg	Val	Thr 40	Ile	Gly	Cys	Gly	Thr 45	Ser	Glu	Asp
20	Ile	Ile 50	Asn	Tyr	Leu	Asn	Trp 55	Tyr	Arg	Lys	Lys	Pro 60	Asp	Gly	Thr	Val
	Glu 65	Leu	Leu	Ile	Tyr	His 70	Thr	Ser	Arg	Leu	Gln 75	Ser	Gly	Val	Pro	Ser 80
25	Arg	Phe	Ser	Gly	Ser 85	Gly	Ser	Gly	Thr	Asp 90	Tyr	Ser	Leu	Thr	Ile 95	Ser
	Asp	Leu	Glu	Gln 100	Glu	Asp	Ile	Ala	Thr 105	Tyr	Phe	Cys	Gln	Gln 110	Gly	Tyr
30	Thr	Leu	Pro 115	Tyr	Thr	Val	Gly	Gly 120	Gly	Thr	Lys	Leu	Glu 125	Ile	Lys	
35	(2)	INFO	RMAT	CION	FOR	SEQ	ID N	10: 3	30:							
40		(i)	(E	L) LE 3) TY 2) SI	ENGTH PE: RANI	I: 41 nucl	.2 ba .eic !SS:	se p acid doub	airs l	3						
		(ii)	MOL		POLC											
45	((iii)	HYE	ОТНЕ	TICA	AL: Y	ES									
		(ix)	•	A) NA	E: ME/K CATI			.1								
50		(ix)) NA	E: AME/F OCATI			-	ide							

		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 3	0:					
5	ATG Met	AAA Lys	TGC Cys 130	AGC Ser	TGG Trp	GTT Val	ATC Ile	TTC Phe 135	TTC Phe	CTG Leu	ATG Met	GCA Ala	GTG Val 140	GTT Val	ACA Thr	GGG	48
		AAT Asn 145														AAG Lys	96
10		GGG Gly															144
15		GAC Asp															192
20		TGG Trp															240
20		AAG Lys															288
25		GCC Ala 225															336
30		TAC Tyr															384
		GGC Gly								G							412
35	(2)	INFO				SEQ CHAR											
40			(<i>I</i>	A) LE 3) Ti	NGTI PE:	H: 13 amir OGY:	7 aπ 10 ac	nino cid									
						PE: ESCRI			SEQ I	D NC): 31	. :					
45	Met 1	Lys	Cys	Ser	Trp 5	Val	Ile	Phe	Phe	Leu 10	Met	Ala	Val	Val	Thr 15	Gly	
	Val	Asn	Ser	Glu 20	Val	Gln	Leu	Gln	Gln 25	Ser	Gly	Ala	Glu	Leu 30	Val	Lys	
50	Pro	Gly	Ala 35	Ser	Val	Asn	Leu	Ser 40	Cys	Thr	Ala	Ser	Gly 45	Phe	Asn	Ile	
	Lys	Asp 50	Thr	Tyr	Met	His	Trp 55	Val	Lys	Gln	Arg	Pro 60	Glu	Gln	Gly	Leu	

	Glu 65	Trp	Ile	Gly	Arg	Ile 70	Asp	Pro	Ala	Asn	Gly 75	Asn	Thr	Lys	Ser	Asp 80	
5	Pro	Lys	Phe	Gln	Ala 85	Lys	Ala	Thr	Ile	Ala 90	Ala	Asp	Thr	Ser	Ser 95	Asn	
	Thr	Ala	Tyr	Leu 100	Gln	Leu	Ser	Ser	Leu 105	Thr	Ser	Glu	Asp	Thr 110	Ala	Val	
10	Tyr	Tyr	Cys 115	Thr	Gly	Gly	Leu	Arg 120	Leu	Arg	Phe	Phe	Asp 125	Tyr	Trp	Gly	
	Gln	Gly 130	Thr	Thr	Leu	Thr	Val 135	Ser	Ser								
15																	
	(2)	INF	ORMA:	rion	FOR	SEQ	ID 1	10: 3	32:								
20		(i)	(I (I	A) LI 3) TY 2) SY	CE CHENGTH PE: TRANI	H: 33 nucl	31 ba Leic ESS:	ase p acio doul	pairs i	5							
25		(ii) (iii)			E TY			A									
30			FE2	ATURI A) NZ		ŒY:	CDS	30									
		(xi)	SEÇ	QUENC	CE DI	ESCRI	PTIC	ON: S	SEQ I	D NO	D: 32	2:					
35		AGA Arg															48
40		CCA Pro 155															96
45		TAC Tyr															144
		ATT															192
50		GGC Gly															240

	GAG Glu	GCT Ala	GAA Glu 220	Asp	GCT Ala	GCC Ala	ACT Thr	TAT Tyr 225	TAC Tyr	TGC Cys	CAG Gln	CAG Gln	TGG Trp 230	AGT Ser	AGT Ser	AAC Asn	288
5	CCA Pro	CCC Pro 235	ATC Ile	ACG Thr	TTC Phe	GGA Gly	GGG Gly 240	GGG Gly	ACC Thr	AAG Lys	CTG Leu	GAA Glu 245	ATA Ile	AAA Lys			330
10	С																331
	(2)	INF	ORMA'	TION	FOR	SEQ	ID 1	. : OV	33:								
15			(1	A) LI B) T	ENGTI YPE :	CHAI H: 1: amir DGY:	10 ar	mino cid									
						YPE: ESCRI	_		SEQ 1	D NO): 33	3:					
20	Ser 1	Arg	Gly	Gln	Ile 5	Val	Leu	Thr	Gln	Ser 10	Pro	Ala	Ile	Met	Ser 15	Ala	
	Ser	Pro	Gly	Glu 20	Lys	Val	Thr	Met	Thr 25	Cys	Ser	Ala	Ser	Ser 30	Ser	Val	
25	Ser	Tyr	Met 35	His	Trp	Tyr	Gln	Gln 40	Lys	Ser	Gly	Thr	Ser 45	Pro	Lys	Arg	
30	Trp	Ile 50	Tyr	Asp	Thr	Ser	Lys 55	Leu	Ala	Ser	Gly	Val 60	Pro	Ala	Arg	Phe	
	Ser 65	Gly	Ser	Gly	Ser	Gly 70	Thr	Ser	Tyr	Ser	Leu 75	Thr	Ile	Ser	Ser	Met 80	
35	Glu	Ala	Glu	Asp	Ala 85	Ala	Thr	Tyr	Tyr	Cys 90	Gln	Gln	Trp	Ser	Ser 95	Asn	
	Pro	Pro	Ile	Thr 100	Phe	Gly	Gly		Thr 105	Lys	Leu	Glu		Lys 110			
40	(2)	INFO	RMAT	ION	FOR	SEQ	ID N	O: 3	4:								
<i>45</i>		(i)	(A (B (C	L) LE () TY () ST	NGTH PE: RAND	ARAC : 5 amin EDNE GY:	amin o ac SS:	o ac id sing	ids								
		(ii)	MOL	ECUL	E TY	PE:	pept	ide									
	(iii)	НҮР	OTHE	TICA	L: Y	ES										
50		(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 34	:					
		Asp 1	Tyr	Gly	Met	Ala 5											

	(2) INFO	RMATION FOR SEQ ID NO: 35:
5	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
10	(ii)	MOLECULE TYPE: peptide
	(iii)	HYPOTHETICAL: YES
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 35:
15	Ala 1	Ile Ser Ser Gly Gly Ser Tyr Ile His Phe Pro Asp Ser Leu Lys 5 10 15
	Gly	
20		
	(2) INFO	RMATION FOR SEQ ID NO: 36:
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 12 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: peptide
30	(iii)	HYPOTHETICAL: YES
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 36:
35	Arg 1	Gly Phe Tyr Gly Asn Tyr Arg Ala Met Asp Tyr 5 10
	(2) INFO	RMATION FOR SEQ ID NO: 37:
40		SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 amino acids
		(B) TYPE: amino acid (C) STRANDEDNESS: single
45		(D) TOPOLOGY: linear
		MOLECULE TYPE: peptide
		HYPOTHETICAL: YES
50		SEQUENCE DESCRIPTION: SEQ ID NO: 37:
	Arg 1	Ala Asn Glu Ser Val Asp His Asn Gly Val Asn Phe Met Asn 5 10 15

	(2) INFORMATION FOR SEQ ID NO: 38:
5	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 7 amino acids(B) TYPE: amino acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
10	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:
15	Ala Ala Ser Asn Gln Gly Ser 1 5
20	(2) INFORMATION FOR SEQ ID NO: 39:
25	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 9 amino acids(B) TYPE: amino acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: peptide
30	(iii) HYPOTHETICAL: YES
-	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:
35	Gln Gln Ser Lys Asp Val Pro Trp Thr 1 5
	(2) INFORMATION FOR SEQ ID NO: 40:
40	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 5 amino acids(B) TYPE: amino acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:
	Ser Tyr Val Ile His 1 5

	(2) INFORMATION FOR SEQ ID NO: 41:
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
10	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:
15	Tyr Ile Asn Pro Tyr Asn Asp Gly Thr Lys Tyr Asn Glu Arg Phe Lys 1 5 10 15
	Gly
20	
	(2) INFORMATION FOR SEQ ID NO: 42:
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 12 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: peptide
30	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:
35	Glu Gly Ile Arg Tyr Tyr Gly Leu Leu Gly Asp Tyr 1 5 10
	(2) INFORMATION FOR SEQ ID NO: 43:
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 11 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:
	Gly Thr Ser Glu Asp Ile Ile Asn Tyr Leu Asn 1 5 10

	(2) INFORMATION FOR SEQ ID NO: 44:
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
10	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
<i>15</i>	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:
	His Thr Ser Arg Leu Gln Ser 1 5
20	(2) INFORMATION FOR SEQ ID NO: 45:
25	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 9 amino acids(B) TYPE: amino acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: peptide
30	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:
35	Gln Gln Gly Tyr Thr Leu Pro Tyr Thr 1 5
	(2) INFORMATION FOR SEQ ID NO: 46:
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 5 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:
	Asp Thr Tyr Met His 1 5

	(2) INFORMATION FOR SEQ ID NO: 47:
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
10	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:
15	Arg Ile Asp Pro Ala Asn Gly Asn Thr Lys Ser Asp Pro Lys Phe Gln 1 5 10 15
	Ala
20	
	(2) INFORMATION FOR SEQ ID NO: 48:
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 9 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: peptide
30	(iii) HYPOTHETICAL: YES
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:
35	Gly Leu Arg Leu Arg Phe Phe Asp Tyr 1 5
	(2) INFORMATION FOR SEQ ID NO: 49:
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 10 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: YES
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:
	Ser Ala Ser Ser Ser Val Ser Tyr Met His 1 5 10

	(2) INFO	DRMATION FOR SEQ ID NO: 50:	
5	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii)	MOLECULE TYPE: peptide	
	(iii)	HYPOTHETICAL: YES	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 50:	
15	Asp 1	Thr Ser Lys Leu Ala Ser 5	
20	(2) INFO	RMATION FOR SEQ ID NO: 51:	
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 10 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: peptide	
		HYPOTHETICAL: YES	
30	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 51:	
35	Gln 1	Gln Trp Ser Ser Asn Pro Pro Ile Thr 5 10	
	(2) INFO	RMATION FOR SEQ ID NO: 52:	
40	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
45	(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
	(iii)	HYPOTHETICAL: YES	
50	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 52:	
- -	GCACCACT	CT CACAGTCTCC TCAGCCAGTA CTAAGGGCC	39

	(2) INFO	RMATION FOR SEQ ID NO: 53:	
5	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
	(iii)	HYPOTHETICAL: YES	
15	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 53:	
	CTTAGTAC	TG GCTGAGGAGA CTGTGAGAGT G	31
20	(2) INFO	RMATION FOR SEQ ID NO: 54:	
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
30	(iii)	HYPOTHETICAL: YES	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 54:	
	GACCAAGT	TG GAAATAAAAC	20
35	(2) INFO	RMATION FOR SEQ ID NO: 55:	
40	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
4 5	(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
	(iii)	HYPOTHETICAL: YES	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 55:	
50	GTACGTTT	TA TTTCCAACTT G	21

	(2) INFORMATION FOR SEQ ID NO: 56:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 97 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:	
15	CAGGAAACAG CTATGACGCG GCCGCCACCA TGGAATGGAG TTGGATATTT CTCTTTCTCC	60
	TGTCAGGAAC TGCAGGTGTC CACTCTGAGG TCCAGCT	97
20	(2) INFORMATION FOR SEQ ID NO: 57:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
25	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
30	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 57:	
	GAATGTGTAT CCAGAAGCCT TGCAGGAAAC CTTCACTGAA GCCCCAGGCT TCTTCACCTC	60
35	AGCTCCAGAC TGCACCAGCT GGACCTCAGA GTGGAC	96
	(2) INFORMATION FOR SEQ ID NO: 58:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 58:	
50	AGGCTTCTGG ATACACATTC ACTAGTTATG TTATTCACTG GGTGCGACAG GCCCCTGGTC	60
	AGGGCCTTGA GTGGATGGGA TATATTAATC CTTACA	96

	(2) INFORMATION FOR SEQ ID NO: 59:	
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 98 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 59:	
15	TGTAGGCTGT GCTCGTGGAC GTGTCTGCAG TGATTGTGAC TCTGCCTTTG AACCTCTCAT	60
	TGTACTTAGT CCCATCATTG TAAGGATTAA TATATCCC	98
20	(2) INFORMATION FOR SEQ ID NO: 60:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 96 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
00	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:	
	GTCCACGAGC ACAGCCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA	60
35	TTACTGTGCG AGAGAAGGAA TTAGGTACTA TGGTCT	96
	(2) INFORMATION FOR SEQ ID NO: 61:	
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 100 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 61:	
50	GTTTTCCCAG TCACGACGGG CCCTTGGTGG AGGCTGAGGA GACTGTGACC AGGGTGCCTT	60
	GGCCCCAGTA GTCTCCCAGT AGACCATAGT ACCTAATTCC	100

	(2)	INF	ORMA	ATION	I FOR	SEC) ID	NO:	62:									
5		(i	((A) I (B) T (C) S	ICE C ENGT YPE: TRAN	H: 4 nuc DEDN	21 b leic ESS:	ase aci dou	pair d	:s								
10		(ii			LE T ESCR							etic	: DNA	."				
		(iii) HY	POTH	ETIC	AL:	YES											
15	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION:1420																	
	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION:157</pre>																	
20		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 6	2:						
	ATG Met	GAA Glu	TGG Trp	AGT Ser	TGG Trp 115	ATA Ile	TTT Phe	CTC Leu	TTT Phe	CTC Leu 120	CTG Leu	TCA Ser	GGA Gly	ACT Thr	GCA Ala 125	GGT Gly		48
25	GTC Val	CAC His	TCT Ser	GAG Glu 130	GTC Val	CAG Gln	CTG Leu	GTG Val	CAG Gln 135	TCT Ser	GGA Gly	GCT Ala	GAG Glu	GTG Val 140	AAG Lys	AAG Lys		96
30	CCT Pro	GGG Gly	GCT Ala 145	TCA Ser	GTG Val	AAG Lys	GTT Val	TCC Ser 150	TGC Cys	AAG Lys	GCT Ala	TCT Ser	GGA Gly 155	TAC Tyr	ACA Thr	TTC Phe	1	L 44
35	ACT Thr	AGT Ser 160	TAT Tyr	GTT Val	ATT Ile	CAC His	TGG Trp 165	GTG Val	CGA Arg	CAG Gln	GCC Ala	CCT Pro 170	GGT Gly	CAG Gln	GGC Gly	CTT Leu	3	192
		TGG Trp															2	240
40		AGG Arg															2	288
45		GCC Ala															3	36
		TAC Tyr															3	84
50		TGG Trp 240											G				4	21

	(2)	INF	ORMA'	rion	FOR	SEQ	ID :	NO:	63:									
5			(1	A) LI B) T	ENGT YPE :		40 a	mino cid	TICS acio									
10						YPE: ESCR			SEQ :	ID NO	D: 63	3:						
	Met 1	Glu	Trp	Ser	Trp 5	Ile	Phe	Leu	Phe	Leu 10	Leu	Ser	Gly	Thr	Ala 15	Gly		
15	Val	His	Ser	Glu 20	Val	Gln	Leu	Val	Gln 25	Ser	Gly	Ala	Glu	Val 30	Lys	Lys		
	Pro	Gly	Ala 35	Ser	Val	Lys	Val	Ser 40	Cys	Lys	Ala	Ser	Gly 45	Tyr	Thr	Phe		
20	Thr	Ser 50	Tyr	Val	Ile	His	Trp 55	Val	Arg	Gln	Ala	Pro 60	Gly	Gln	Gly	Leu		
	Glu 65	Trp	Met	Gly	Tyr	Ile 70	Asn	Pro	Tyr	Asn	Asp 75	Gly	Thr	Lys	Tyr	Asn 80		
25	Glu	Arg	Phe	Lys	Gly 85	Arg	Val	Thr	Ile	Thr 90	Ala	Asp	Thr	Ser	Thr 95	Ser		
	Thr	Ala	Tyr	Met 100	Glu	Leu	Ser	Ser	Leu 105	Arg	Ser	Glu	Asp	Thr 110	Ala	Val		
30	Tyr	Tyr	Cys 115	Ala	Arg	Glu	Gly	Ile 120	Arg	Tyr	Tyr	Gly	Leu 125	Leu	Gly	Asp		
	Tyr	Trp 130	Gly	Gln	Gly	Thr	Leu 135	Val	Thr	Val	Ser	Ser 140						
35	(2)	INFC	RMAT	NOI	FOR	SEQ	ID N	10: 6	54:									
40		(i)	(A (E (C	L) LE 3) TY 1) SI	NGTE PE: RANI	IARAC I: 87 nucl EDNE	bas eic SS:	se pa ació sing	irs I									
45		(ii)							clei sc =			tic	DNA"					
	((iii) (xi)						N: 5	EQ I	D NO): 64	•						
50	CAGO												GTTC	CT 1	GGTC	TCCT	}	60
50		CTCTG																87

	(2) INFORMATION FOR SEQ ID NO: 65:	
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 83 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 65:	
15	GATGGTGACT CTGTCTCCTA CAGAGGCAGA CAGGGAGGAT GGAGACTGTG TCATCTGGAT	60
	ATCACATCTG ATGTCTTGAA AAC	83
20	(2) INFORMATION FOR SEQ ID NO: 66:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 92 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
25	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 66:	
	TAGGAGACAG AGTCACCATC ACTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60
35	GGTATCAACA GAAACCAGGG AAAGCCCCTA AG	92
	(2) INFORMATION FOR SEQ ID NO: 67:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 90 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 67:	
50	TTCCAGACCC GCTGCCACTG AACCTTGATG GGACTCCTGA CTGTAATCTT GATGTGTGGT	60
	AGATCAGGAG CTTAGGGGCT TTCCCTGGTT	90

	(2) INFORMATION FOR SEQ ID NO: 68:	
5	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 68:	
15	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTAGTCTGC AACCTGAAGA	60
	TTTTGCCACT TACTACTGCC AACAGGGT	88
22	(2) INFORMATION FOR SEQ ID NO: 69:	
20	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 91 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
25	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 69:	
	GTTTTCCCAG TCACGACCGT ACGTTTATT TCCACCTTGG TCCCTTGGCC GAACGTGTAC	60
35	GGAAGCGTAT AACCCTGTTG GCAGTAGTAA G	91
	(2) INFORMATION FOR SEQ ID NO: 70:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 382 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear 	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
50	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION:1381	
	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION:160</pre>	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 70:																		
5		ATG Met														CAA Gln		48
10		ATC Ile																96
70		TCT Ser																144
15		ATC Ile 190																192
20		CTC Leu																240
		TTC Phe																288
25		CTG Leu																336
30		CTT Leu																381
	С																	382
35	(2)	INF	(i) (i) (i) (i) (ii) (ii) (ii) (ii) (ii	SEQUI A) LI 3) T	ENCE ENGTI	CHAP I: 12 amir	RACTE 27 an	ERIST	rics:									
40			MOI	LECUI	DPOLO	PE:	prot	ein	SEQ 1	ID NO): 7 <u>1</u>	.:						
45	Met 1	Met	Ser	Ser	Ala 5	Gln	Phe	Leu	Gly	Leu 10	Leu	Leu	Leu	Cys	Phe 15	Gln		
	Asp	Ile	Arg	Cys 20	Asp	Ile	Gln	Met	Thr 25	Gln	Ser	Pro	Ser	Ser 30	Leu	Ser		
50	Ala	Ser	Val 35	Gly	Asp	Arg	Val	Thr 40	Ile	Thr	Cys	Gly	Thr 45	Ser	Glu	Asp		
	Ile	Ile 50	Asn	Tyr	Leu	Asn	Trp 55	Tyr	Gln	Gln	Lys	Pro 60	Gly	Lys	Ala	Pro		

	Lys Leu 65	Leu Ile	Tyr His 70	Thr Ser	Arg	Leu	Gln 75	Ser	Gly	Val	Pro	Ser 80			
5	Arg Phe	Ser Gly	Ser Gly 85	Ser Gly	Thr	Asp 90	Phe	Thr	Leu	Thr	Ile 95	Ser			
	Ser Leu	Gln Pro (100	Glu Asp	Phe Ala	Thr 105	Tyr	Tyr	Cys	Gln	Gln 110	Gly	Tyr			
10		Pro Tyr '	Thr Phe	Gly Gln 120	Gly	Thr	Lys	Val	Glu 125	Ile	Lys				
	(2) INFORMATION FOR SEQ ID NO: 72:														
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear														
20															
	(iii) HYPOTHETICAL: YES														
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 72: GTCCACGAGC ACAGCCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA														
	GTCCACGAGC ACAGCCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA														
	TCTCTGTGGG AGAGAAGGAA TTAGGTACTA TGGTCT														
30	(2) INFORMATION FOR SEQ ID NO: 73:														
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 421 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear														
	(ii)	MOLECULE	TYPE: 0					+ia	וו מזאת						
40	(iii)	нүротнет			-	.	iiciie	CIC	DNA						
45	(ix)		ME/KEY: (CATION:1												
40	(ix)		ME/KEY: SCATION:1		ide										
50	(xi)	SEQUENCE	DESCRI	PTION: S	SEQ I	D NO	: 73	:							
	ATG GAA '												48		

	GTC Val	CAC His	Ser	GAG Glu	GTC Val	CAG Gln	CTG Leu 150	GTG Val	CAG Gln	TCT Ser	GGA Gly	GCT Ala 155	GAG Glu	GTG Val	AAG Lys	AAG Lys		96
5		Gly			GTG Val											TTC Phe 175		144
10	ACT Thr	AGT Ser	TAT Tyr	GTT Val	ATT Ile 180	CAC His	TGG Trp	GTG Val	CGA Arg	CAG Gln 185	GCC Ala	CCT Pro	GGT Gly	CAG Gln	GGC Gly 190	CTT Leu		192
					TAT Tyr													240
15					GGC Gly													288
20					GAG Glu													336
25					AGA Arg													384
					GGC Gly 260								G					421
30	(2)	INF	ORMAT	rion	FOR	SEQ	ID N	io: 7	74:									
35	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 140 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear																	
-					E TY		_		EQ I	D NO): 74	l:						
40	Met 1	Glu	Trp	Ser	Trp 5	Ile	Phe	Leu	Phe	Leu 10	Leu	Ser	Gly	Thr	Ala 15	Gly		
	Val	His	Ser	Glu 20	Val	Gln	Leu	Val	Gln 25	Ser	Gly	Ala	Glu	Val 30	Lys	Lys		
45	Pro	Gly	Ala 35	Ser	Val	Lys	Val	Ser 40	Cys	Lys	Ala	Ser	Gly 45	Tyr	Thr	Phe		
	Thr	Ser 50	Tyr	Val	Ile	His	Trp 55	Val	Arg	Gln	Ala	Pro 60	Gly	Gln	Gly	Leu		
50	Glu 65	Trp	Met	Gly	Tyr	Ile 70	Asn	Pro	Tyr	Asn	Asp 75	Gly	Thr	Lys	Tyr	Asn 80		
	Glu	Arg	Phe	Lys	Gly 85	Arg	Val	Thr	Ile	Thr 90	Ala	Asp	Thr	Ser	Thr 95	Ser		

	Thr Ala Tyr Met Glu Leu Ser Ser Leu Arg Ser Glu Asp Thr Ala Val 100 105 110														
5	Tyr Leu Cys Gly Arg Glu Gly Ile Arg Tyr Tyr Gly Leu Leu Gly Asp 115 120 125														
	Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Ser 130 135 140														
10	(2) INFORMATION FOR SEQ ID NO: 75:														
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear														
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>														
20	(iii) HYPOTHETICAL: YES (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 75:														
	AGGCCTTCC GTCGATCCCA TATATTAATC CTTACA														
25	AGGGCCTTGC GTGGATGGGA TATATTAATC CTTACA														
	(2) INFORMATION FOR SEQ ID NO: 76:														
30	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 98 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 														
35	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>														
	(iii) HYPOTHETICAL: YES														
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 76:														
40	TGTAGGCTGT GCTCGTGGAC CTGTCTGCAG TGATTGTGAC TCTGCCTTTG AACCTCTCAT	60													
	TGTACTTAGT CCCATCATTG TAAGGATTAA TATATCCC	98													
45	(2) INFORMATION FOR SEQ ID NO: 77:														
50	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 421 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear														
	(ii) MOLECULE TYPE: other nucleic acid														

			(A) D	ESCR	IPTI	ON:	/d	esc	= "s	ynth	etic	DNA	n			
		(iii) HY	POTH	ETIC.	AL:	YES										
5		(ix	(.	ATUR: A) N. B) L	AME/			20									
10		(ix	(.	ATURI A) Ni B) L	AME/				tide								
		(xi) SE	QUEN	CE D	ESCR:	IPTI	ON:	SEQ	ID N	0: 7	7:					
		GAA				_											48
15	Met	Glu	Trp	Ser	145	He	Pne	Leu	Pne	150	Leu	Ser	GIĄ	Thr	A1a 155	GIA	
		CAC															96
	vai	His	ser	160	vai	GIII	Leu	Val	165	ser	GIY	AIA	GIU	170	гув	гÀг	
20		GGG Gly						Ser					Gly				144
								180					185				
25		AGT Ser 190															192
		TGG															240
30	Ala 205	Trp	Met	Gly	Tyr	11e 210	Asn	Pro	Tyr	Asn	Asp 215	Gly	Thr	Lys	Tyr	Asn 220	
30		AGG															288
	Glu	Arg	Pne	гÀг	225	Arg	vai	Tnr	lle	230	Ala	Asp	Arg	ser	235	ser	
35		GCC															336
	Inr	Ala	TŸĽ	240	GIU	Leu	ser	ser	245	AIG	ser	GIU	Asp	250	Ala	Val	
		CTC Leu															384
40	171	Deu	255	Cly	9	Olu	Cly	260	9	-7-	-7-	017	265	204	O17	p	
		TGG Trp											G				421
	ıyı	270	O17	U 111	O ₂ y	****	275	· u _		,	501	280					
45	(2)	INF	ORMA:	rion	FOR	SEQ	ID 1	40: '	78:								
				SEQUI													
			(1	3) T 3) T	YPE:	amir	no a	cid		_							
50		(ii)		LECUI													
		(xi)	SE	QUEN	CE DI	ESCR:	IPTI	ON: S	SEQ :	ID N	D: 78	3:					

	Met 1	Glu	Trp	Ser	Trp 5	Ile	Phe	Leu	Phe	Leu 10	Leu	Ser	Gly	Thr	Ala 15	Gly	
5	Val	His	Ser	Glu 20	Val	Gln	Leu	Val	Gln 25	Ser	Gly	Ala	Glu	Val 30	Lys	Lys	
	Pro	Gly	Ala 35	Ser	Val	Lys	Val	Ser 40	Cys	Lys	Ala	Ser	Gly 45	Tyr	Thr	Phe	
10	Thr	Ser 50	Tyr	Val	Ile	His	Trp 55	Val	Arg	Gln	Ala	Pro 60	Gly	Gln	Gly	Leu	
	Ala 65	Trp	Met	Gly	Tyr	Ile 70	Asn	Pro	Tyr	Asn	Asp 75	Gly	Thr	Lys	Tyr	Asn 80	
15	Glu	Arg	Phe	Lys	Gly 85	Arg	Val	Thr	Ile	Thr 90	Ala	Asp	Arg	Ser	Thr 95	Ser	
	Thr	Ala	Tyr	Met 100	Glu	Leu	Ser	Ser	Leu 105	Arg	Ser	Glu	Asp	Thr 110	Ala	Val	
20	Tyr	Leu	Cys 115	Gly	Arg	Glu	Gly	Ile 120	Arg	Tyr	Tyr	Gly	Leu 125	Leu	Gly	Asp	
	Tyr	Trp 130	Gly	Gln	Gly	Thr	Leu 135	Val	Thr	Val	Ser	Ser 140					
25	(2)	INFO	ORMAT	NOI	FOR	SEQ	ID N	10: 7	9:								
		(i)		UENC													
30	(A) LENGTH: 96 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear																
	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"																
35	(iii) HYPOTHETICAL: YES																
		(xi)	SEÇ	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	: 79	:					
	AGGC	TTCT	rgg A	TACA	CATI	'C AC	TAGT	TATG	TTA	TTCA	.CTG	GGTG	CGAC	CAG A	AGGCC	TGGTC	60
40	AGGG	CCTI	GC G	TGGA	TGGG	А ТА	TATI	AATC	CTI	'ACA							96
	(2)	INFO	RMAT	CION	FOR	SEQ	ID N	10: 8	0:								
45		(i)	(F (E	QUENC LE B) TY C) ST D) TO	NGTH PE: RAND	: 98 nucl	bas eic SS:	e pa acid sing	irs								
50		(ii)		ECUL								tic	DNA"	ı			
	(iii)	HYE	POTHE	TICA	L: Y	ES										

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 80:	
	TGTAGACTGT GCTCGTGGAC CTGTCTGAAG TGATTGTGAC TTTGCCTTTG AACCTCTCAT	60
5	TGTACTTAGT CCCATCATTG TAAGGATTAA TATATCCC	98
	(2) INFORMATION FOR SEQ ID NO: 81:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
15	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 81:	
20	GTCCACGAGC ACAGTCTACA TGGAGCTCAG TTCGCTGAGA TCTGAGGACA CGGCGGTGTA	60
	TCTCTGTGGG AGAGAAGGAA TTAGGTACTA TGGTCT	96
25	(2) INFORMATION FOR SEQ ID NO: 82:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 421 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: double	
30	(D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
35	(iii) HYPOTHETICAL: YES	
	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION:1420	
40	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION:157</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 82:	
45	ATG GAA TGG AGT TGG ATA TTT CTC TTT CTC CTG TCA GGA ACT GCA GGT Met Glu Trp Ser Trp Ile Phe Leu Phe Leu Leu Ser Gly Thr Ala Gly 145 150 155	48
	GTC CAC TCT GAG GTC CAG CTG GTG CAG TCT GGA GCT GAG GTG AAG AAG Val His Ser Glu Val Gln Leu Val Gln Ser Gly Ala Glu Val Lys Lys 160 165 170	96
50	CCT GGG GCT TCA GTG AAG GTT TCC TGC AAG GCT TCT GGA TAC ACA TTC Pro Gly Ala Ser Val Lys Val Ser Cys Lys Ala Ser Gly Tyr Thr Phe	144

		175		180	185	
5		Tyr Val I			AGG CCT GGT CA Arg Pro Gly Gl 200	
10				Pro Tyr Asn	GAT GGG ACT AM Asp Gly Thr Ly 215	
		Phe Lys G			TCA GAC AGG TO Ser Asp Arg Se	
15					TCT GAG GAC AC Ser Glu Asp Th 25	ar Ala Val
20					TAT GGT CTA CT Tyr Gly Leu Le 265	
		Gly Gln G		GTC ACA GTC Val Thr Val		421
25	(2) INF	ORMATION F				
30	,,,	(A) LENGER (B) TYPE (D) TOPE	GTH: 140 ar E: amino ac OLOGY: line	nino acids cid ear		
) MOLECULE) SEQUENCE	_	cein ON: SEQ ID NO	: 83:	
35	Met Glu 1	Trp Ser T	rp Ile Phe 5	Leu Phe Leu 10	Leu Ser Gly Th	r Ala Gly 15
	Val His	Ser Glu V	al Gln Leu	Val Gln Ser 25	Gly Ala Glu Va 3	l Lys Lys O
40	Pro Gly	Ala Ser V	al Lys Val	Ser Cys Lys	Ala Ser Gly Ty 45	r Thr Phe
	Thr Ser		le His Trp 55	Val Arg Gln	Arg Pro Gly Gl 60	n Gly Leu
45	Ala Trp 65	Met Gly T	yr Ile Asn 70	Pro Tyr Asn	Asp Gly Thr Ly 75	s Tyr Asn 80
	Glu Arg	•	ly Lys Val 85	Thr Ile Thr 90	Ser Asp Arg Se	r Thr Ser 95
50	Thr Val	Tyr Met G 100	lu Leu Ser	Ser Leu Arg 105	Ser Glu Asp Th 11	
	Tyr Leu	Cys Gly A	rg Glu Gly	Ile Arg Tyr 120	Tyr Gly Leu Le 125	eu Gly Asp

	Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Ser 130 135 140	
5	(2) INFORMATION FOR SEQ ID NO: 84:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 92 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
15	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 84:	
	TAGGAGACAG AGTCACCATC ACTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60
20	GGTATCGGCA GAAACCAGGG AAAGCCCCTG AA	92
	(2) INFORMATION FOR SEQ ID NO: 85:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 90 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
30	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 85:	
	TTCCAGACCC GCTGCCACTG AACCTTGATG GGACTCCTGA CTGTAATCTT GATGTGTGGT	60
	AGATCAGGAG TTCAGGGGCT TTCCCTGGTT	90
40	(2) INFORMATION FOR SEQ ID NO: 86:	
45	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 91 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
50	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 86:	

	GTTTTCCCAG TCACGACCGT ACGTTTTATT TCCACCTTGG TCCCTTGGCC GACCGTGTAC														
	GGAAGCGTAT AACCCTGTTG GCAGTAGTAA G	91													
5	(2) INFORMATION FOR SEQ ID NO: 87:														
10	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 382 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear 														
15	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"														
	(iii) HYPOTHETICAL: YES (ix) FEATURE:														
20	(A) NAME/KEY: CDS (B) LOCATION:1381 (ix) FEATURE:														
	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION:160</pre>														
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 87:														
	ATG ATG TCC TCT GCT CAG TTC CTT GGT CTC CTG TTG CTC TGT TTT CAA Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln 145 150 155	48													
30	GAT ATC AGA TGT GAT ATC CAG ATG ACA CAG TCT CCA TCC TCC CTG TCT Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser 160 165 170	96													
35	GCC TCT GTA GGA GAC AGA GTC ACC ATC ACT TGC GGG ACA AGT GAG GAC Ala Ser Val Gly Asp Arg Val Thr Ile Thr Cys Gly Thr Ser Glu Asp 175 180 185	L44													
	ATT ATC AAT TAT TTA AAC TGG TAT CGG CAG AAA CCA GGG AAA GCC CCT Ile Ile Asn Tyr Leu Asn Trp Tyr Arg Gln Lys Pro Gly Lys Ala Pro 190 195 200	L92													
40	GAA CTC CTG ATC TAC CAC ACA TCA AGA TTA CAG TCA GGA GTC CCA TCA Glu Leu Leu Ile Tyr His Thr Ser Arg Leu Gln Ser Gly Val Pro Ser 205 210 215 220	240													
45	AGG TTC AGT GGC AGC GGG TCT GGA ACA GAT TTC ACT CTC ACC ATT AGT Arg Phe Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser 225 230 235	288													
50	AGT CTG CAA CCT GAA GAT TTT GCC ACT TAC TAC TGC CAA CAG GGT TAT Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Gly Tyr 240 245 250	336													
	ACG CTT CCG TAC ACG GTC GGC CAA GGG ACC AAG GTG GAA ATA AAA Thr Leu Pro Tyr Thr Val Gly Gln Gly Thr Lys Val Glu Ile Lys 255 260 265	381													

	С	382
5	(2) INFORMATION FOR SEQ ID NO: 88:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 127 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 88:	
15	Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln 1 5 10 15	
	Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser 20 25 30	
20	Ala Ser Val Gly Asp Arg Val Thr Ile Thr Cys Gly Thr Ser Glu Asp 35 40 45	
	Ile Ile Asn Tyr Leu Asn Trp Tyr Arg Gln Lys Pro Gly Lys Ala Pro 50 55 60	
25	Glu Leu Leu Ile Tyr His Thr Ser Arg Leu Gln Ser Gly Val Pro Ser 65 70 75 80	
	Arg Phe Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser 85 90 95	
30	Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln Gly Tyr 100 105 110	
	Thr Leu Pro Tyr Thr Val Gly Gln Gly Thr Lys Val Glu Ile Lys 115 120 125	
35	(2) INFORMATION FOR SEQ ID NO: 89:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 92 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
45	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 89:	
50	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT GGTATCGGCA GAAACCAGGG AAAGCCCCTG AA	60 92

	(2) INFORMATION FOR SEQ ID NO: 90:														
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear														
10	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>														
	(iii) HYPOTHETICAL: YES														
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 90:														
15	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTGACCTGC AACCTGAAGA														
	TTTTGCCACT TACTACTGCC AACAGGGT														
20	(2) INFORMATION FOR SEQ ID NO: 91: (i) SEQUENCE CHARACTERISTICS:														
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 382 base pairs														
	(B) TYPE: nucleic acid (C) STRANDEDNESS: double														
25	(D) TOPOLOGY: linear														
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>														
	(iii) HYPOTHETICAL: YES														
30	(ix) FEATURE:														
	(A) NAME/KEY: CDS (B) LOCATION:1381														
35	(ix) FEATURE:														
	(A) NAME/KEY: sig_peptide (B) LOCATION:160														
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 91:														
40	ATG ATG TCC TCT GCT CAG TTC CTT GGT CTC CTG TTG CTC TGT TTT CAA Met Met Ser Ser Ala Gln Phe Leu Gly Leu Leu Leu Cys Phe Gln	48													
	130 135 140														
	GAT ATC AGA TGT GAT ATC CAG ATG ACA CAG TCT CCA TCC TCC CTG TCT Asp Ile Arg Cys Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser	96													
45	145 150 155														
	GCC TCT GTA GGA GAC AGA GTC ACC ATC GGT TGC GGG ACA AGT GAG GAC Ala Ser Val Gly Asp Arg Val Thr Ile Gly Cys Gly Thr Ser Glu Asp	44													
	160 165 170 175														
50	ATT ATC AAT TAT TTA AAC TGG TAT CGG CAG AAA CCA GGG AAA GCC CCT Ile Ile Asn Tyr Leu Asn Trp Tyr Arg Gln Lys Pro Gly Lys Ala Pro	92													
	180 185 190														

		CTC Leu															:	240
5		TTC Phe															:	288
10		CTG Leu 225															:	336
15		CTT Leu															:	381
	С																:	382
20	(2)	INFO																
25		,	(I	1) LI 3) TY	ENGTI PE :	H: 12 amir	RACTI 27 and 10 ac line	nino cid										
			MOI SEÇ				_		SEQ]	D NO): 92	2:						
30	Met 1	Met	Ser	Ser	Ala 5	Gln	Phe	Leu	Gly	Leu 10	Leu	Leu	Leu	Cys	Phe 15	Gln		
	Asp	Ile	Arg	Cys 20	Asp	Ile	Gln	Met	Thr 25	Gln	Ser	Pro	Ser	Ser 30	Leu	Ser		
35	Ala	Ser	Val 35	Gly	Asp	Arg	Val	Thr 40	Ile	Gly	Cys	Gly	Thr 45	Ser	Glu	Asp		
40	Ile	Ile 50	Asn	Tyr	Leu	Asn	Trp 55	Tyr	Arg	Gln	Lys	Pro 60	Gly	Lys	Ala	Pro		
	Glu 65	Leu	Leu	Ile	Tyr	His 70	Thr	Ser	Arg	Leu	Gln 75	Ser	Gly	Val	Pro	Ser 80		
45	Arg	Phe	Ser	Gly	Ser 85	Gly	Ser	Gly	Thr	Asp 90	Phe	Thr	Leu	Thr	Ile 95	Ser		
	Asp	Leu	Gln	Pro 100	Glu	Asp	Phe	Ala	Thr 105	Tyr	Tyr	Cys	Gln	Gln 110	Gly	Tyr		
50	Thr	Leu	Pro 115	Tyr	Thr	Val	Gly	Gln 120	Gly	Thr	Lys	Val	Glu 125	Ile	Lys			

	(2) INFORMATION FOR SEQ ID NO: 93:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 83 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 93:	
15	GATGGTGACT CTGTCTCCTA CAGAGGCAGA CAGGGAGGAT GTAGCCTGTG TCATCTGGAT	60
	ATCACATCTG ATGTCTTGAA AAC	83
20	(2) INFORMATION FOR SEQ ID NO: 94:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 92 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"	
30	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 94:	
	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60
35	GGTATCGGAA GAAACCAGGG AAAGCCCCTG AA	92
	(2) INFORMATION FOR SEQ ID NO: 95:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 95:	
50	CAGTGGCAGC GGGTCTGGAA CAGATTTCAC TCTCACCATT AGTGACCTGC AACCTGAAGA	60
	TTTTGCCACT TACTTTTGCC AACAGGGT	88

	(2) I	NFOR	TAMS	ION	FOR	SEQ	ID N	iO: 9	6:								
5		(i)	(A) (B) (C)	LE: TY	NGTH PE :	: 91 nucl EDNE	bas eic SS:	STIC e pa acid sing	irs								
10	(ii)							clei sc =			tic	DNA"				
	(i	ii)	HYPO	OTHE	TICA	L: Y	ES										
15	(xi)	SEQ	JENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	: 96	:					
	GTTTT	CCCA	G T	CACG	ACCG	T AC	GTTI	TATI	TCC	ACCI	TGG	TCCC	TTGG	CC G	ACCG	TGTAC	60
	GGAAG	CGTA	AT A	ACCC	TGTT	G GC	'AAAA	GTAA	G								91
20	(2) I	NFOR	TAMS	ION	FOR	SEQ	ID N	iO: 9	7:								
25		(i)	(A (B (C	LE TY	NGTH PE :	: 38 nucl EDNE	2 ba eic SS:	acid doub	airs l	i							
30			(A) DE		PTIC	N:		clei sc =			etic	DNA"				
<i>35</i>	·	ix)	(A (B FEA') NA) LO TURE) NA	ME/K CATI	ON:1	38 sig_	pept	ide								
40	,	/# \	•-						SEQ I	וח אור)· 97	7 •					
	ATG A												CTC	ጥርያጥ	ԴԴ	۵۵	48
45	Met M	iet s	Ser 130	Ser	Ala	Gln	Phe	Leu 135	Gly	Leu	Leu	Leu	Leu 140	Cys	Phe	Gln	10
	GAT A Asp I	ATC A lle A	AGA Arg	TGT Cys	GAT Asp	ATC Ile	CAG Gln 150	ATG Met	ACA Thr	CAG Gln	GCT Ala	ACA Thr 155	TCC Ser	TCC Ser	CTG Leu	TCT Ser	96
50	GCC T Ala S 160	CT (Ser \	GTA Val	GGA Gly	GAC Asp	AGA Arg 165	GTC Val	ACC Thr	ATC Ile	GGT Gly	TGC Cys 170	GGG Gly	ACA Thr	AGT Ser	GAG Glu	GAC Asp 175	144

5	ATT ATO															192
5	GAA CTC Glu Leu															240
10	AGG TTO															288
15	GAC CTC Asp Leu 225	Gln														336
	ACG CTT Thr Lev 240															381
20	С															382
	(2) INF	ORMAT	NOI	FOR	SEQ	ID N	10: 9	8:								
25		(B	LE () TY	ENGTI (PE :	CHAF H: 12 amir OGY:	27 an	nino cid									
30) MOL						SEQ 1	D NO): 98	B:					
	Met Met 1	Ser	Ser	Ala 5	Gln	Phe	Leu	Gly	Leu 10	Leu	Leu	Leu	Cys	Phe 15	Gln	
35	Asp Ile	Arg	Cys 20	Asp	Ile	Gln	Met	Thr 25	Gln	Ala	Thr	Ser	Ser 30	Leu	Ser	
	Ala Ser	Val 35	Gly	Asp	Arg	Val	Thr 40	Ile	Gly	Cys	Gly	Thr 45	Ser	Glu	Asp	
40	Ile Ile	Asn	Tyr	Leu	Asn	Trp 55	Tyr	Arg	Lys	Lys	Pro 60	Gly	Lys	Ala	Pro	
	50					33										
	Glu Leu 65)	Ile	Tyr	His 70		Ser	Arg	Leu	Gln 75	Ser	Gly	Val	Pro	Ser 80	
45	Glu Lev	Leu			70	Thr				75					80	
45 50	Glu Leu 65	Leu Ser	Gly Pro 100	Ser 85 Glu	70 Gly Asp	Thr Ser	Gly Ala	Thr Thr 105	Asp 90 Tyr	75 Phe Phe	Thr Cys	Leu Gln	Thr Gln 110	Ile 95 Gly	80 Ser	

	(2) INFORMATION FOR SEQ ID NO: 99:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 92 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic DNA"</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 99:	
15	TAGGAGACAG AGTCACCATC GGTTGCGGGA CAAGTGAGGA CATTATCAAT TATTTAAACT	60
	GGTATCGGAA GAAACCAGGG AAAGCCGTTG AA	92
20	(2) INFORMATION FOR SEQ ID NO: 100:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 90 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
30	(iii) HYPOTHETICAL: YES	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 100:	
	TTCCAGACCC GCTGCCACTG AACCTTGATG GGACTCCTGA CTGTAATCTT GATGTGTGT	60
35	AGATCAGGAG TTCAACGGCT TTCCCTGGTT	90
	(2) INFORMATION FOR SEQ ID NO: 101:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear 	
45	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 101:	
50	CAGTGGCAGC GGGTCTGGAA CAGATTATAC TCTCACCATT AGTGACCTGC AACCTGAAGA	60
	TTTTGCCACT TACTTTTGCC AACAGGGT	88

	(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:	102:								
5		(i)	() () ()	QUENCA) LIB) TO	engti YPE : IRANI	H: 3: nuc: DEDNI	32 ba leic ESS:	ase p acio doul	pair:	S							
10		(ii)		LECUI A) Di								etic	DNA				
		(iii)	HY:	POTH	ETIC	AL: 3	YES										
15		(ix)	(2	ATURI A) NI B) L(AME/I			31									
20	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION:160 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 102:</pre>																
		(xi)	SE	QUEN	CE DI	ESCR:	[PTIC	ON: S	SEQ :	ID NO): 10	02:					
25		ATG Met															48
		ATC Ile 145															96
30		TCT Ser															144
35		ATC Ile											_			_	192
40		CTC Leu															240
		TTC Phe															288
45		CTG Leu 225															336
50		CTT Leu															381
	С																382

	(2)	INFO	RMAI	NOI	FOR	SEQ	ID I	10: 1	103:								
5		((<i>P</i>	L) LE		I: 12 amir	27 ar										
					E TY		_	cein ON: S	SEQ :	D NO): 10	3:					
10	Met N	Met	Ser	Ser	Ala 5	Gln	Phe	Leu	Gly	Leu 10	Leu	Leu	Leu	Cys	Phe 15	Gln	
	Asp :	Ile	Arg	Cys 20	Asp	Ile	Gln	Met	Thr 25	Gln	Ala	Thr	Ser	Ser 30	Leu	Ser	
15	Alas	Ser	Val 35	Gly	Asp	Arg	Val	Thr 40	Ile	Gly	Cys	Gly	Thr 45	Ser	Glu	Asp	
	Ile	Ile 50	Asn	Tyr	Leu	Asn	Trp 55	Tyr	Arg	Lys	Lys	Pro 60	Gly	Lys	Ala	Val	
20	Glu 1 65	Leu	Leu	Ile	Tyr	His 70	Thr	Ser	Arg	Leu	Gln 75	Ser	Gly	Val	Pro	Ser 80	
05	Arg 1	Phe	Ser	Gly	Ser 85	Gly	Ser	Gly	Thr	Asp 90	Tyr	Thr	Leu	Thr	Ile 95	Ser	
25	Asp 1	Leu	Gln	Pro 100	Glu	Asp	Phe	Ala	Thr 105	Tyr	Phe	Cys	Gln	Gln 110	Gly	Tyr	
30	Thr 1		Pro 115	Tyr	Thr	Val	Gly	Gln 120	Gly	Thr	Lys	Val	Glu 125	Ile	Lys		
	(2)	INFO	RMAT	rion	FOR	SEQ	ID 1	MO: 3	104:								
35		(i)	(<i>I</i> (E	A) LI 3) T C) S	engti Pe :	nuci DEDNI	5 ba: leic ESS:	ISTIC se pa acio sing ear	airs d								
40		(ii)						er n /d				etic	DNA	11			
	(iii)	НУІ	POTH	ETIC	AL:	YES										
45	GCTT							ON:	SEQ	ID N	0: 1	04:					25
	(2)	INFO	ORMA'	rion	FOR	SEQ	ID	NO:	105:								
50			SE(QUEN A) L B) T C) S	CE C ENGT YPE:	HARA H: 3 nuc DEDN	CTER 0 ba leic ESS:	ISTI se p aci sin	CS: airs d								

	(11)	MOLECU (A) I	ESCRI						id nthe	tic 1	'' A'					
5	(iii)	нүротн	ETICA	L: Y	ES											
	(xi)	SEQUEN	ICE DE	SCRI	PTIO	N: S1	EQ II	OM C	: 10	5:						
	AAGGATCC	TG GCAC	TCATT	T AC	CCAG	AGAC										30
10	(2) INFO	RMATION	FOR	SEQ :	ID NO	D: 10	06:									
15	(i)	(B) T	CE CH ENGTH YPE: TRAND	: 31: amino EDNE:	3 am: o ac: SS: s	ino a id sing!	acids	3								
	(ii) MOLECULE TYPE: peptide															
20	(iii) HYPOTHETICAL: YES															
	(xi)	SEQUEN	CE DE	SCRI	PTIO	N: SI	EQ II	ONO	: 106	5:						
	Asp 1	Leu Le	u Pro	Asp 5	Glu	Lys	Ile	Ser	Leu 10	Leu	Pro	Pro	Val	Asn 15	Phe	
25	Thr	Ile Ly	s Val 20	Thr	Gly	Leu	Ala	Gln 25	Val	Leu	Leu	Gln	Trp 30	Lys	Pro	
	Asn	Pro As	-	Glu	Gln	Arg	Asn 40	Val	Asn	Leu	Glu	Tyr 45	Gln	Val	Lys	
30	Ile	Asn Al	a Pro	Lys	Glu	Asp 55	Asp	Tyr	Glu	Thr	Arg 60	Ile	Thr	Glu	Ser	
35	Lys 65	Cys Va	l Thr	Ile	Leu 70	His	Lys	Gly	Phe	Ser 75	Ala	Ser	Val	Arg	Thr 80	
33	Ile	Leu Gl	n Asn	Asp 85	His	Ser	Leu	Leu	Ala 90	Ser	Ser	Trp	Ala	Ser 95	Ala	
40	Glu	Leu Hi	s Ala 100		Pro	Gly	Ser	Pro 105	Gly	Thr	Ser	Val	Val 110	Asn	Leu	
		Cys Th												Arg	Ser	
45	Tyr	Gln Va	l Ser	Leu	His	Cys 135	Thr	Trp	Leu	Val	Gly 140	Thr	Asp	Ala	Pro	
	Glu 145	Asp Th	ır Gln	Tyr	Phe 150	Leu	Tyr	Tyr	Arg	Tyr 155	Gly	Ser	Trp	Thr	Glu 160	
50	Glu	Cys G	n Glu	Tyr 165	Ser	Lys	Asp	Thr	Leu 170	Gly	Arg	Asn	Ile	Ala 175	Cys	
	Trp	Phe Pi	o Arg 180		Phe	Ile	Leu	Ser 185	Lys	Gly	Arg	Asp	Trp 190	Leu	Ala	

	Val	Leu	Val 195	Asn	Gly	Ser	Ser	Lys 200	His	Ser	Ala	Ile	Arg 205	Pro	Phe	Asp
5	Gln	Leu 210	Phe	Ala	Leu	His	Ala 215	Ile	Asp	Gln	Ile	Asn 220	Pro	Pro	Leu	Asn
10	Val 225	Thr	Ala	Glu	Ile	Glu 230	Gly	Thr	Arg	Leu	Ser 235	Ile	Gln	Trp	Glu	Lys 240
	Pro	Val	Ser	Ala	Phe 245	Pro	Ile	His	Cys	Phe 250	Asp	Tyr	Glu	Val	Lys 255	Ile
15	His	Asn	Thr	Arg 260	Asn	Gly	Tyr	Leu	Gln 265	Ile	Glu	Lys	Leu	Met 270	Thr	Asn
	Ala	Phe	Ile 275	Ser	Ile	Ile	Asp	Asp 280	Leu	Ser	Lys	Tyr	Asp 285	Val	Gln	Val
20	Arg	Ala 290	Ala	Val	Ser	Ser	Met 295	Cys	Arg	Glu	Ala	Gly 300	Leu	Trp	Ser	Glu
25	Trp 305	Ser	Gln	Pro	Ile	Tyr 310	Val	Gly	Lys							

30 Claims

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- 1. An antibody which reacts specifically with a human interleukin-5 receptor α chain.
- 2. The antibody of claim 1, which is selected from monoclonal antibodies, humanized antibodies, single chain antibodies and disulfide-stabilized antibodies.
 - 3. The antibody of claim 2, which is a monoclonal antibody.
- 40 **4.** The monoclonal antibody of claim 3, which binds to an epitope at 1-313 positions from the N-terminal amino acid of a human interleukin-5 receptor α chain and which reacts specifically with the human interleukin-5 receptor α chain by immunocyte staininig.
- 5. The monoclonal antibody of claim 3, which binds to an epitope at 1-313 positions from the N-terminal amino acid of the human interleukin-5 receptor α chain and which inhibits a biological activity of human interleukin-5.
 - 6. The monoclonal antibody of claim 4, which belongs to IgG1 subclass, CDR sequences in the variable region (V region) of the heavy chain (H chain) of the antibody being the following amino acid sequences:

CDR1: Asp Tyr Gly Met Ala

CDR2: Ala IIe Ser Ser Gly Gly Ser Tyr IIe His Phe Pro Asp Ser Leu Lys Gly CDR3: Arg Gly Phe Tyr Gly Asn Tyr Arg Ala Met Asp Tyr

and CDR sequences in the V region of the light chain (L chain) being the following amino acid sequences:

CDR1: Arg Ala Asn Glu Ser Val Asp His Asn Gly Val Asn Phe Met Asn

CDR2: Ala Ala Ser Asn Gln Gly Ser

CDR3: Gln Gln Ser Lys Asp Val Pro Trp Thr.

- The monoclonal antibody of claim 6, which is monoclonal antibody KM1257 produced by hybridoma KM1257 (FERM BP-5133).
- 8. The monoclonal antibody of claim 5, which belongs to IgG1 subclass, CDR sequences in the V region of the H chain of the antibody being the following amino acid sequences:

CDR1: Ser Tyr Val IIe His

CDR2: Tyr lle Asn Pro Tyr Asn Asp Gly Thr Lys Tyr Asn Glu Arg Phe Lys Gly

CDR3: Glu Gly lle Arg Tyr Tyr Gly Leu Leu Gly Asp Tyr

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and CDR sequences in the V region of the L chain being the following amino acid sequences:

CDR1: Gly Thr Ser Glu Asp lle lle Asn Tyr Leu Asn

CDR2: His Thr Ser Arg Leu Gln Ser

CDR3: Gln Gln Gly Tyr Thr Leu Pro Tyr Thr.

- The monoclonal antibody of claim 8, which is monoclonal antibody KM1259 produced by hybridoma KM1259 (FERM BP-5134).
- 20 10. The monoclonal antibody of claim 5, which belongs to IgG1 subclass, CDR sequences in the V region of the H chain of the antibody being the following amino acid sequences:

CDR1: Asp Thr Tyr Met His

CDR2: Arg Ile Asp Pro Ala Asn Gly Asn Thr Lys Ser Asp Pro Lys Phe Gln Ala

CDR3: Gly Leu Arg Leu Arg Phe Phe Asp Tyr

and CDR sequences in the V region of the L chain being the following amino acid sequences:

CDR1: Ser Ala Ser Ser Ser Val Ser Tyr Met His

CDR2: Asp Thr Ser Lys Leu Ala Ser

CDR3: Gln Gln Trp Ser Ser Asn Pro Pro lle Thr

- **11.** The monoclonal antibody of claim 10, which is monoclonal antibody KM1486 produced by hybridoma KM1486 (FERM BP-5651).
- 12. Hybridoma KM1257 (FERM BP-5133) which produces the monoclonal antibody of claim 6.
- 13. Hybridoma KM1259 (FERM BP-5134) which produces the monoclonal antibody of claim 8.
- 40 14. Hybridoma KM1486 (FERM BP-5651) which produces the monoclonal antibody of claim 10.
 - 15. The antibody of claim 2, which is a humanized antibody.
 - 16. The humanized antibody of claim 15, which recognizes an epitope at 1-313 positions from the N-terminal amino acid of a human interleukin-5 receptor α chain and which reacts specifically with the human interleukin-5 receptor α chain by immunocyte staining.
 - **17.** The humanized antibody of claim 15, which recognizes an epitope at 1-313 positions from the N-terminal amino acid of a human interleukin-5 receptor *α* chain and which inhibits a biological activity of human interleukin-5.
 - 18. The antibody of claim 16 or 17, which belongs to human antibody IgG class.
 - 19. The antibody of claim 15, 16 or 17, which is a human chimeric antibody.
- 20. The antibody of claim 19, wherein the human chimeric antibody is a chimeric antibody comprising the V region of the H chain and the V region of the L chain of a non-human animal antibody, as well as the constant region (C region) of the H chain and the C region of the L chain of a human antibody.
 - 21. The antibody of claim 20, wherein the V region of the H chain of the antibody comprises the amino acid sequence

of SEQ ID NO: 24 and the V region of the L chain of the antibody comprises the amino acid sequence of SEQ ID NO: 25.

- 22. The antibody of claim 20, wherein the V region of the H chain of the antibody comprises the amino acid sequence of SEQ ID NO: 26 and the V region of the L chain of the antibody comprises the amino acid sequence of SEQ ID NO: 27.
 - 23. The antibody of claim 21 being antibody KM1399, wherein the C region of the H chain of the antibody is in a human antibody IgG1 subclass.
 - 24. The antibody of claim 21 being antibody KM7399, wherein the C region of the H chain of the antibody is in a human antibody IgG4 subclass.
 - 25. Transformant KM1399 (FERM BP-5650) which produces the antibody of claim 23.
 - 26. Transformant KM7399 (FERM BP-5649) which produces the antibody of claim 24.
 - 27. The antibody of claim 15, wherein the humanized antibody is a human CDR-grafted antibody.
- 28. The antibody of claim 17, wherein the human CDR-grafted antibody is obtained by replacing CDR sequences in the V region of the H chain and the V region of the L chain of a human antibody with CDR sequences in the V region of the H chain and the V region of the L chain of a non-human animal antibody.
- 29. The antibody of claim 28, wherein a CDR sequence in the V region of the H chain of the antibody comprises a CDR sequence in the V region of the H chain of the antibody of claim 8 and a CDR sequence in the V region of the L chain of the antibody comprises a CDR sequence in the V region of the L chain of the antibody of claim 8.
 - 30. The antibody of claim 28, wherein a CDR sequence in the V region of the H chain of the antibody comprises a CDR sequence in the V region of the H chain of the antibody of claim 10 and a CDR sequence in the V region of the L chain of the antibody comprises a CDR sequence in the V region of the L chain of the antibody of claim 10.
 - **31.** The antibody of claim 29 being antibody KM8399, wherein the C region of the H chain of the antibody belongs to human antibody IgG1 subclass.
- 35 32. The antibody of claim 29 being antibody KM9399, wherein the C region of the H chain of the antibody is in a human antibody IgG4 subclass.
 - 33. Transformant KM8399 (FERM BP-5648) which produces the antibody of claim 31.
- 40 34. Transformant KM9399 (FERM BP-5647) which produces the antibody of claim 32.
 - 35. Tandem cassette vector pKANTEX93.

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- **36.** A process of producing a humanized antibody by means of a transformant containing tandem cassette vector, pKANTEX93.
 - 37. The antibody of claim 2, wherein the antibody is a single chain antibody.
- **38.** The single chain antibody of claim 37, which recognizes an epitope at 1-313 positions from the N-terminal amino acid of a human interleukin-5 receptor α chain and which inhibits a biological activity of human interleukin-5.
 - **39.** The antibody of claim 38, wherein the single chain antibody comprises the V region of the H chain and the V region of the L chain of a humanized antibody.
- 40. The single chain antibody of claim 38, wherein CDR sequences in the V region of the H chain and the V region of the L chain comprise CDR sequences in the V region of the H chain and the V region of the L chain of the monoclonal antibody of claim 8.
 - 41. The single chain antibody of claim 38, wherein CDR sequences in the V region of the H chain and the V region of

the L chain comprise CDR sequences in the V region of the H chain and the V region of the L chain of the monoclonal antibody of claim 8.

- 42. The antibody of claim 2, which is a disulfide-stabilized antibody.
- **43.** The disulfide-stabilized antibody of claim 42, which recognizes an epitope at 1-313 positions from the N-terminal amino acid of a human interleukin-5 receptor α chain and which inhibits a biological activity of human interleukin-5.
- 44. The disulfide-stabilized antibody of claim 43, which comprises the V region of the H chain and the V region of the L chain of a humanized antibody.
 - **45.** The disulfide-stabilized antibody of claim 43, wherein CDR sequences in the V region of the H chain and the V region of the L chain comprise CDR sequences in the V region of the H chain and the V region of the L chain of the monoclonal antibody of claim 8.
 - **46.** The single chain antibody of claim 43, wherein CDR sequences in the V region of the H chain and the V region of the L chain comprise CDR sequences in the V region of the H chain and the V region of the L chain of the monoclonal antibody of claim 10.
- 47. A peptide comprising a CDR sequence in the V region of the H chain or the V region of the L chain of the monoclonal antibody of claim 6, 8 or 10 which has a reactivity with a human interleukin-5 receptor α chain.
 - **48.** A human interleukin-5 receptor α chain antigen comprising a protein which can be obtained by transfecting into a host cell a vector capable of expressing an entire region or a partial fragment of a human interleukin-5 receptor α chain either as such or as a fusion protein, expressing in culture, isolating and purifying the protein or fragment thereof.
 - **49.** The human interleukin-5 receptor α chain antigen of claim 48, which has the amino acid sequence of SEQ ID NO: 91.
 - **50.** A method for detecting a human interleukin-5 receptor α chain immunologically by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
- 51. A method for immunological detection of a cell expressing a human interleukin-5 receptor α chain on the surface by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
 - **52.** A method for detecting human eosinophils immunologically by means ox any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
- 40 53. A method for detecting and determining a soluble human interleukin-5 receptor α chain immunologically by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
 - 54. A method for preventing eosinophilia by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
- 55. A method for treating allergic diseases such as chronic bronchial asthma by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
 - **56.** A method for treating eosinophilia-involving diseases by means of any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46.
 - **57.** An eosinophilia-inhibiting agent comprising any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46 as an active ingredient.
- 58. A therapeutic agent for allergic diseases comprising any one of the antibodies of claims 1-11, 15-24, 27-32 and 37-46 as an active ingredient.

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FIG.1

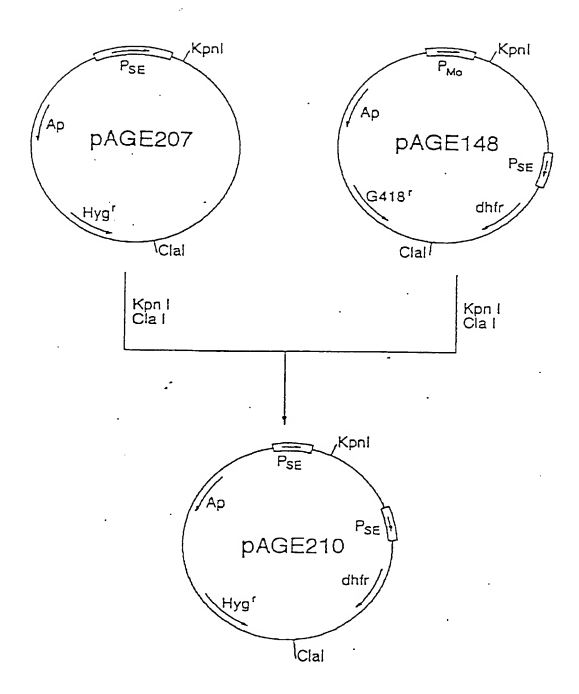


FIG.2

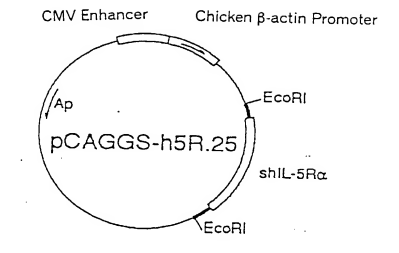


FIG.3

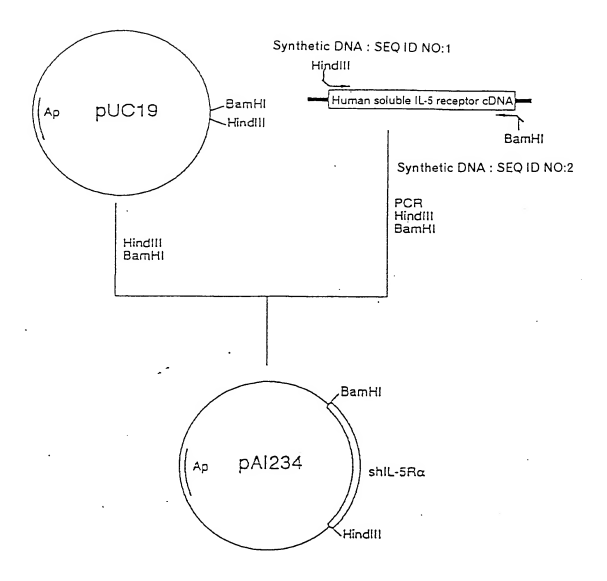


FIG.4

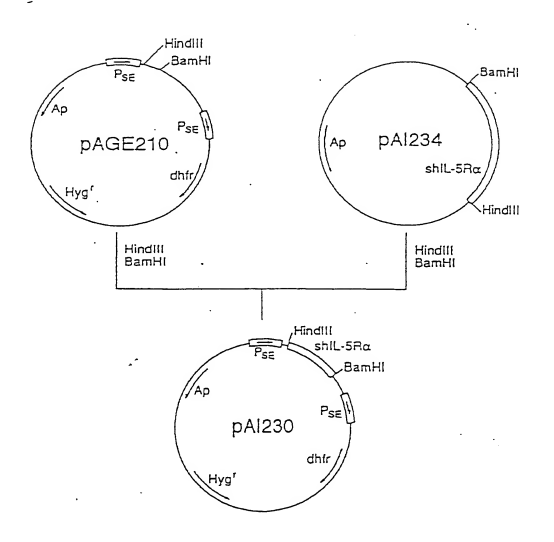


FIG.5

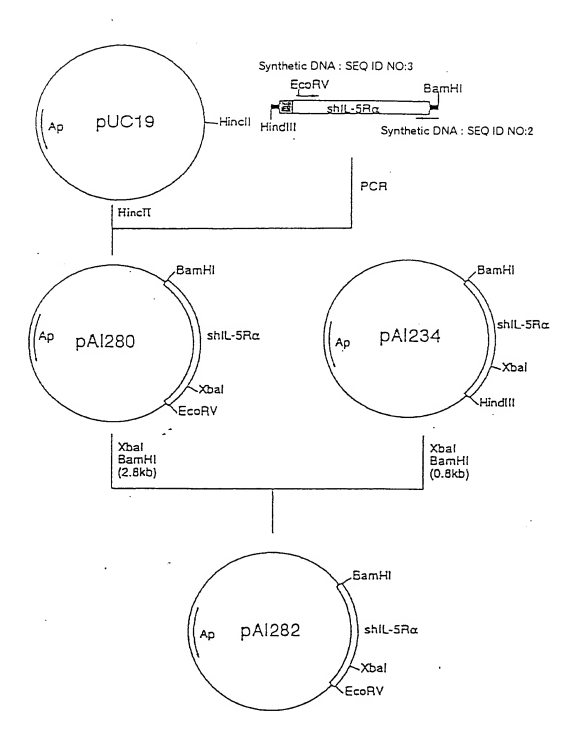


FIG.6

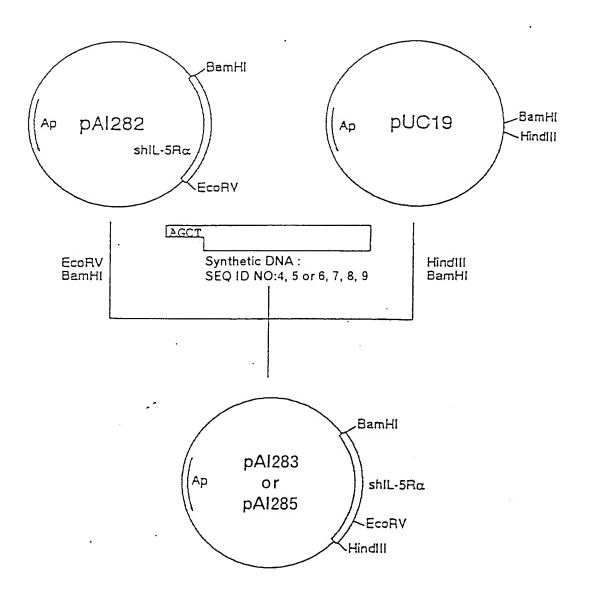


FIG.7

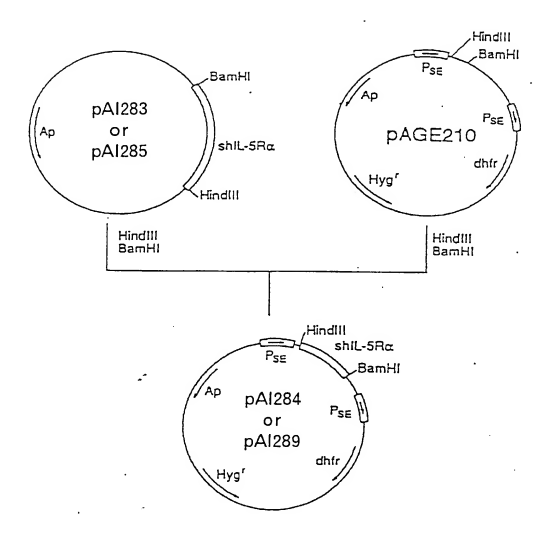


FIG.8

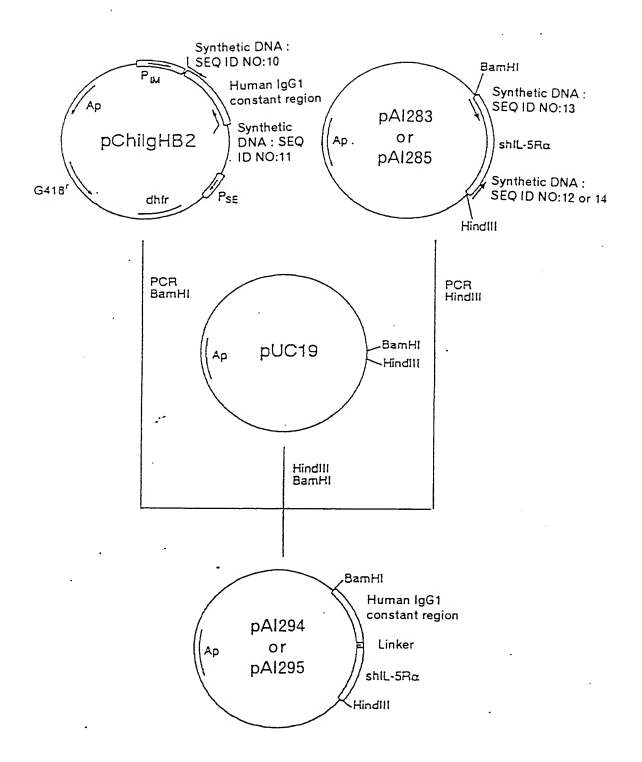


FIG.9

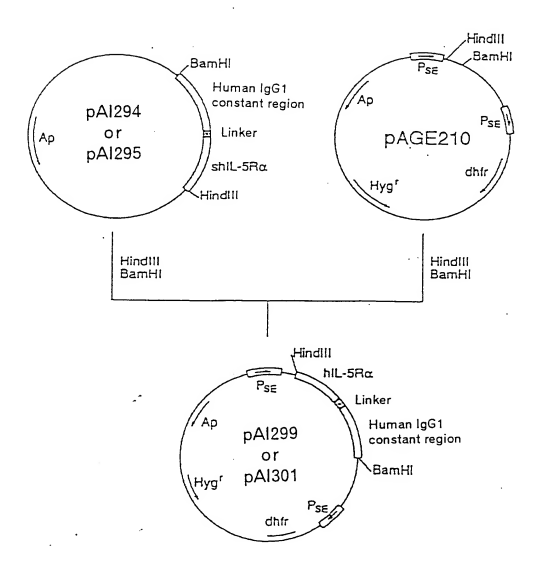


FIG.10

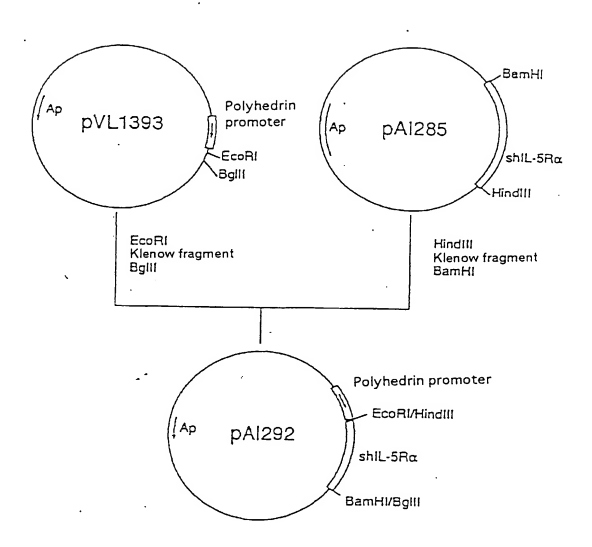


FIG.11

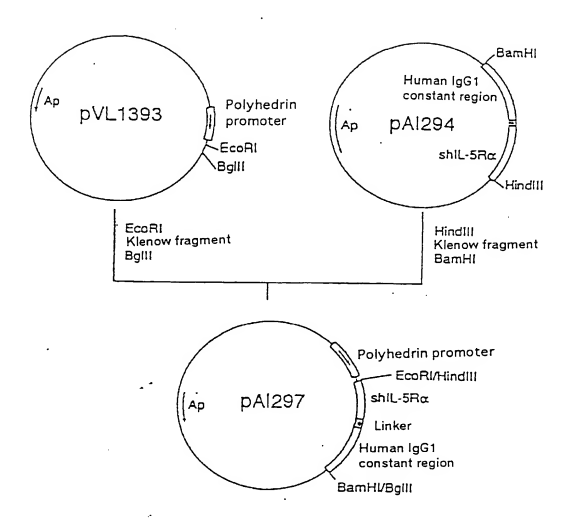


FIG.12

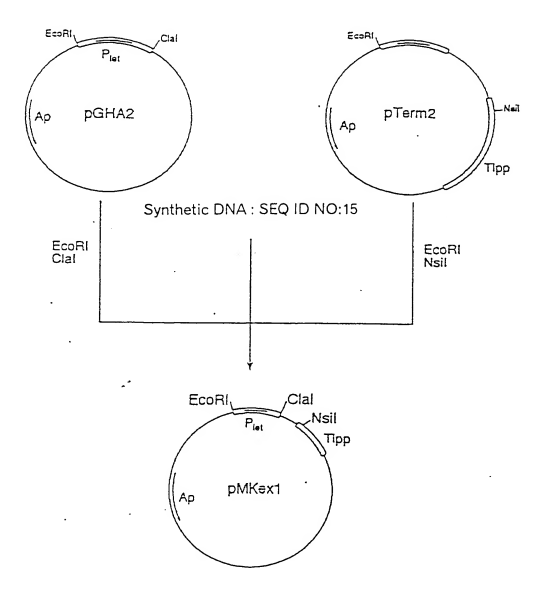


FIG.13

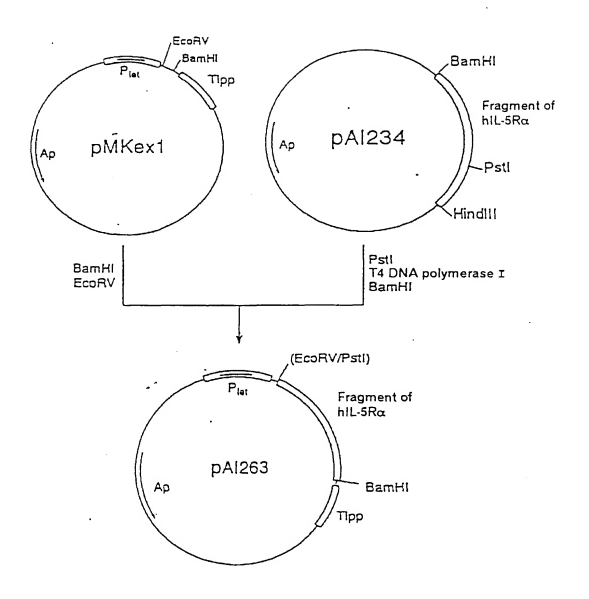


FIG.14

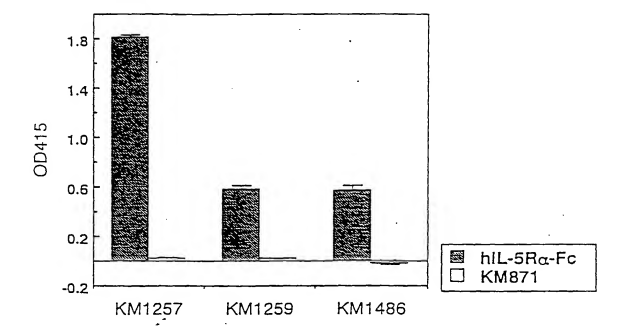


FIG.15

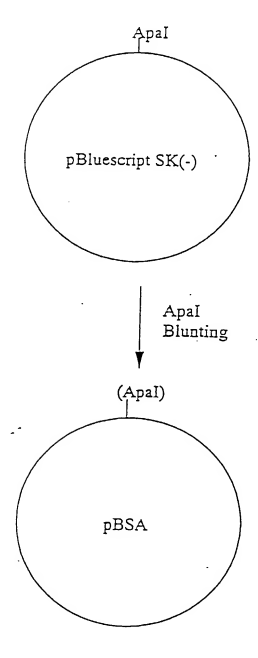


FIG.16

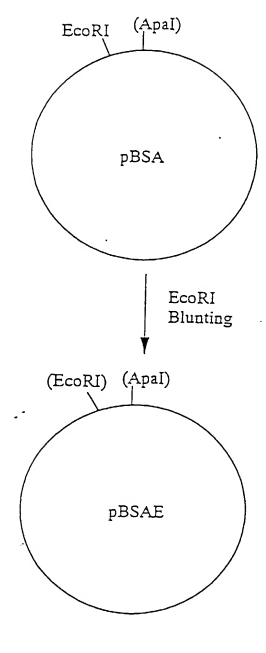


FIG.17

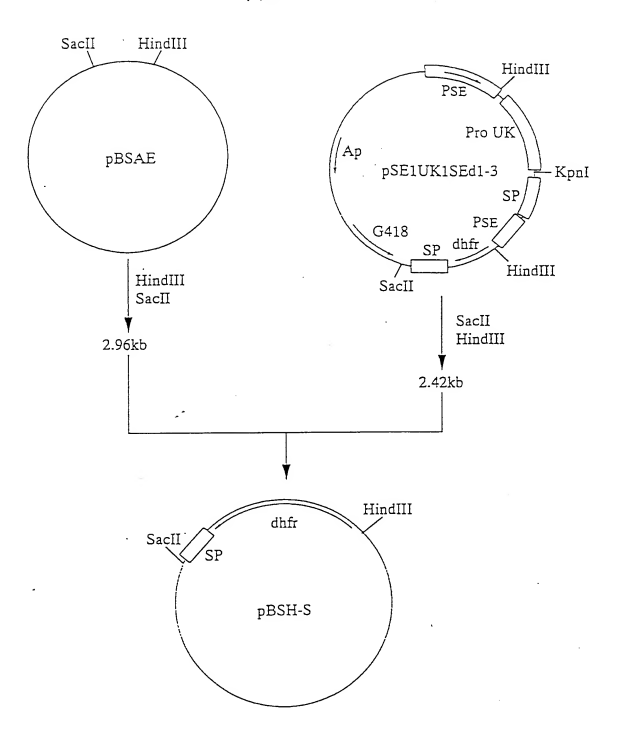


FIG.18

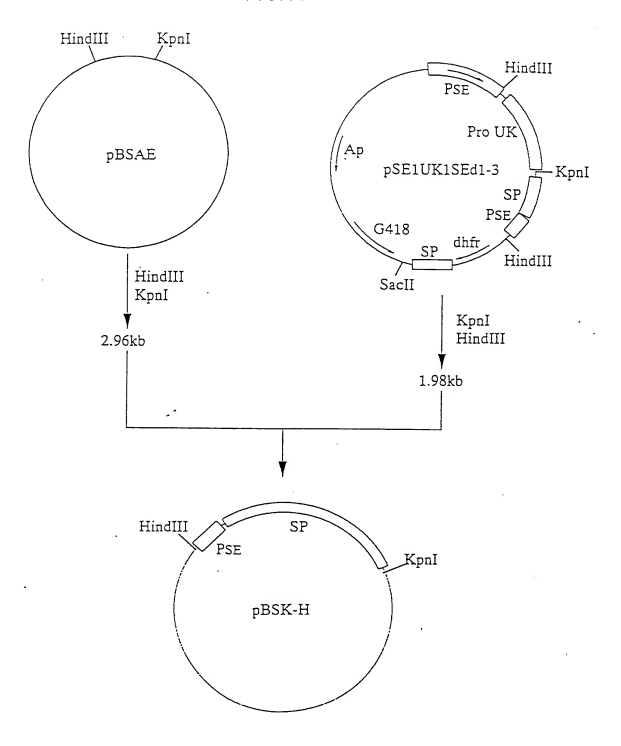


FIG.19

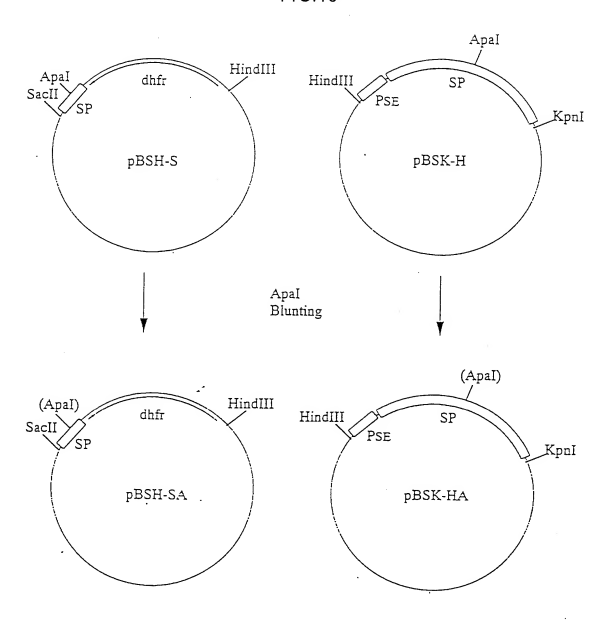


FIG.20

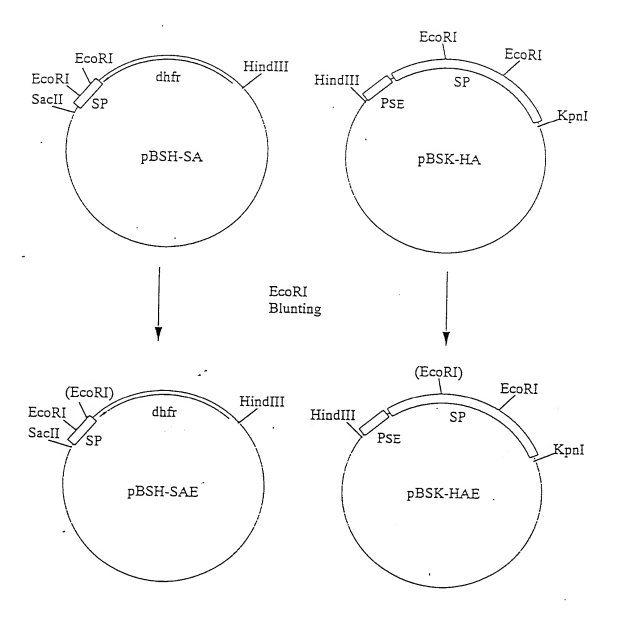


FIG.21

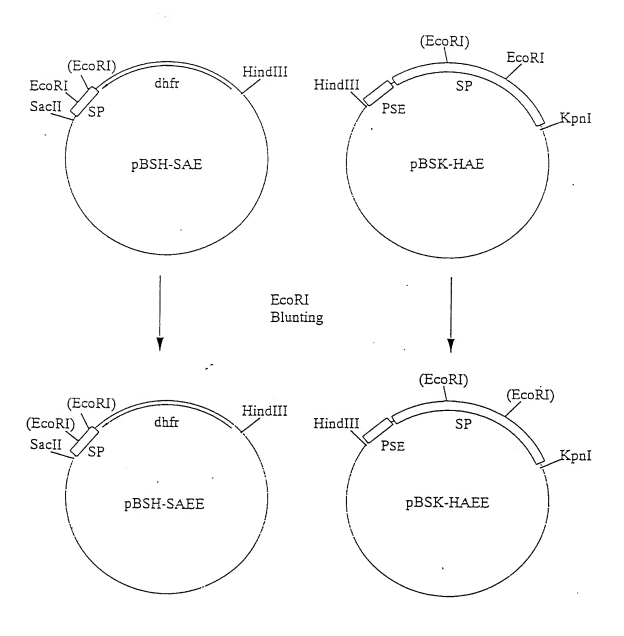


FIG.22

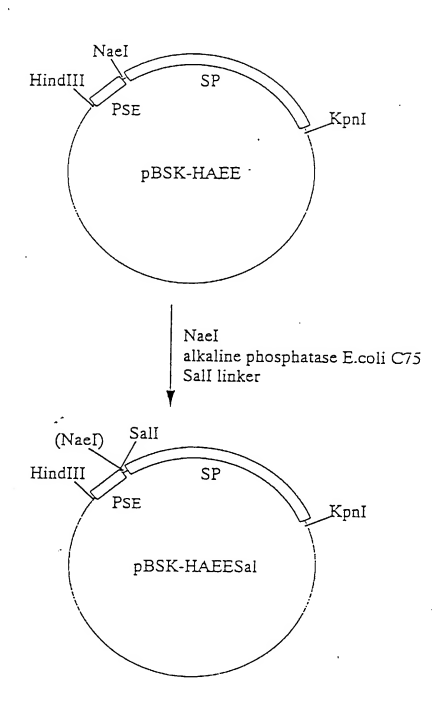


FIG.23

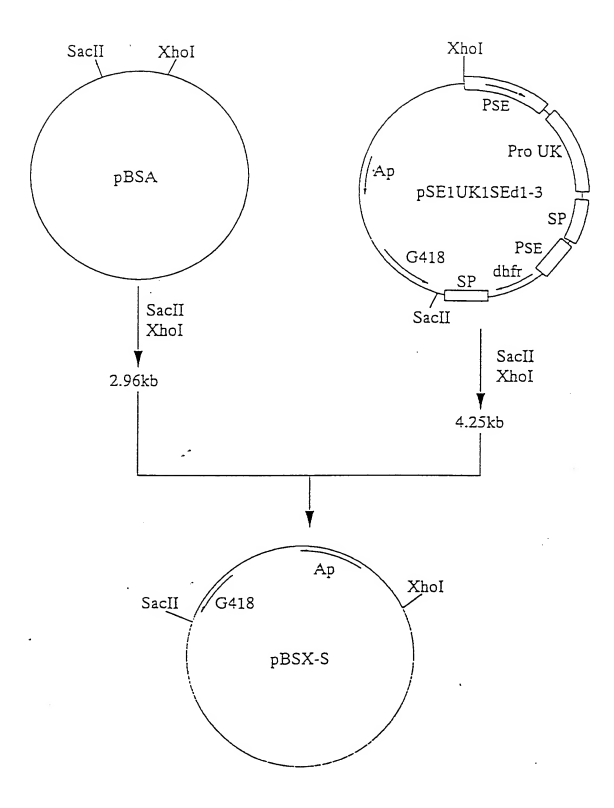


FIG.24

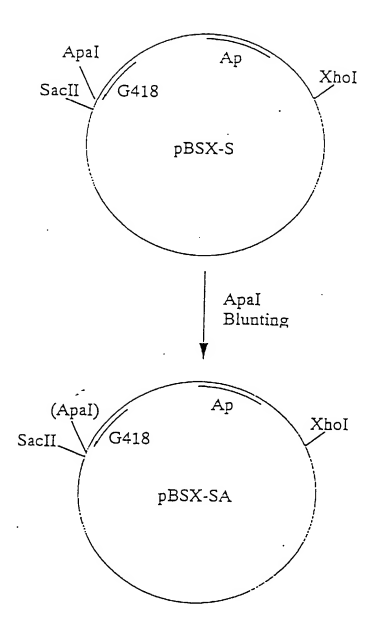


FIG.25

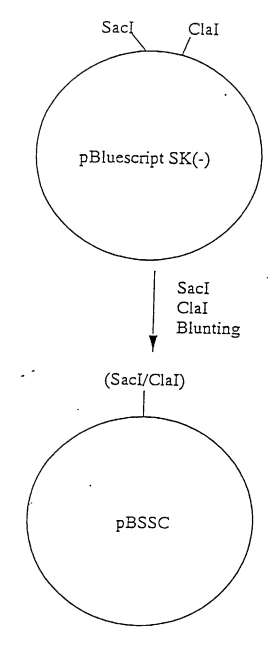


FIG.26

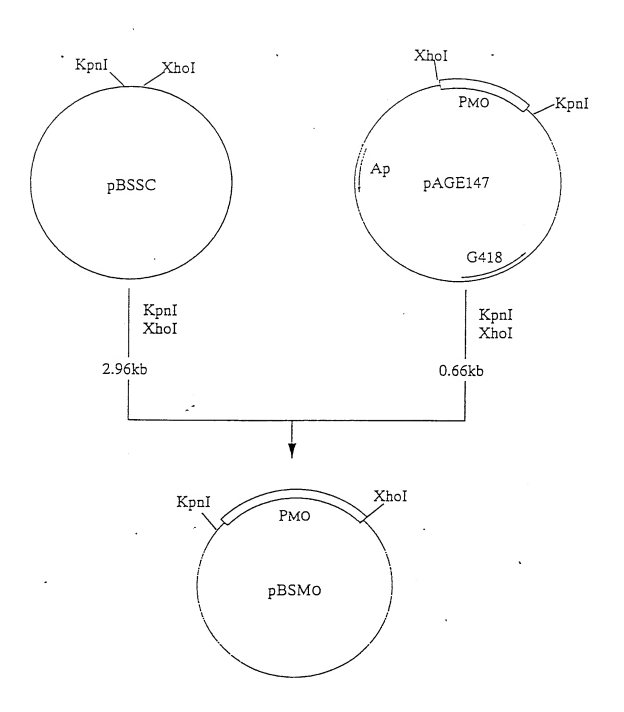


FIG.27

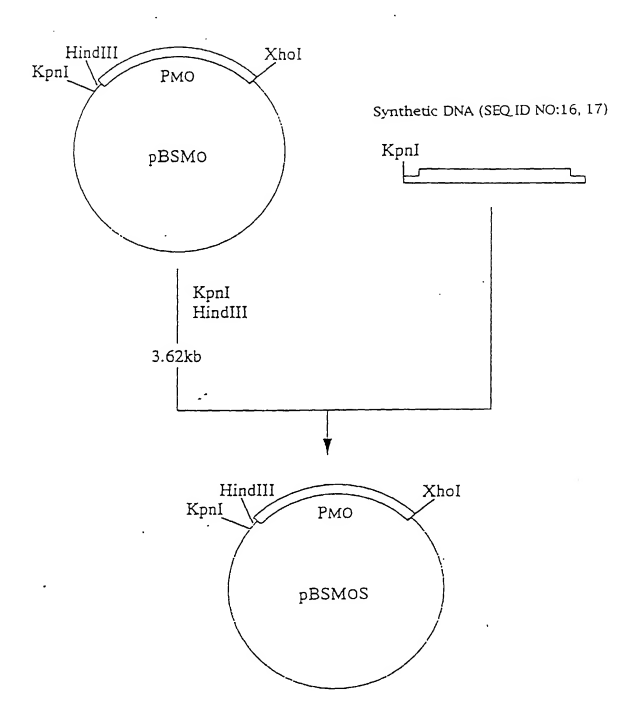


FIG.28

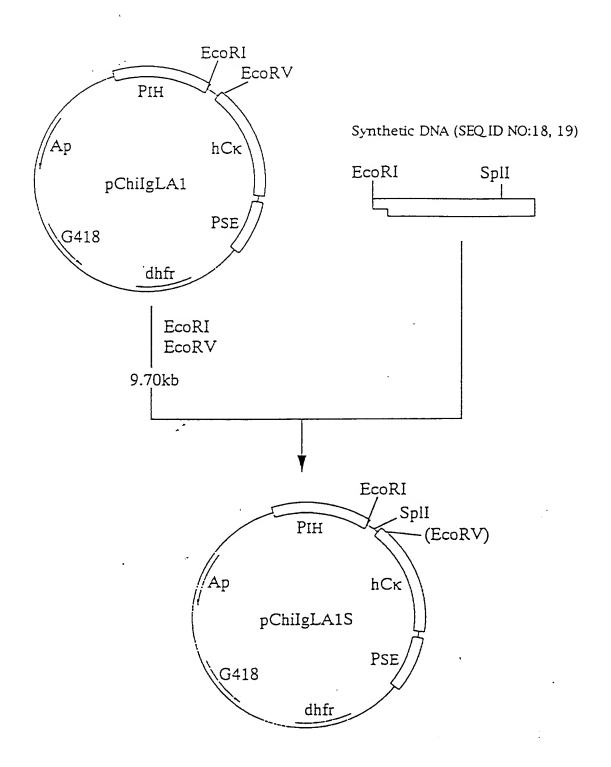


FIG.29

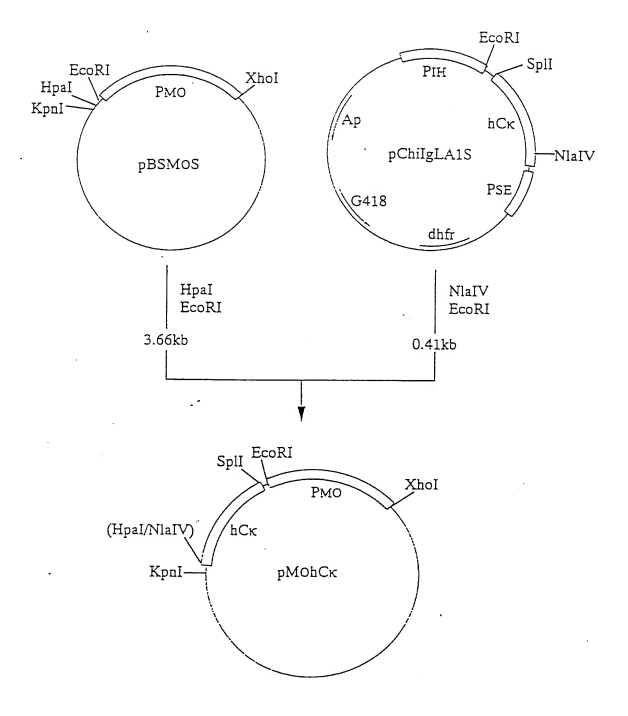


FIG.30

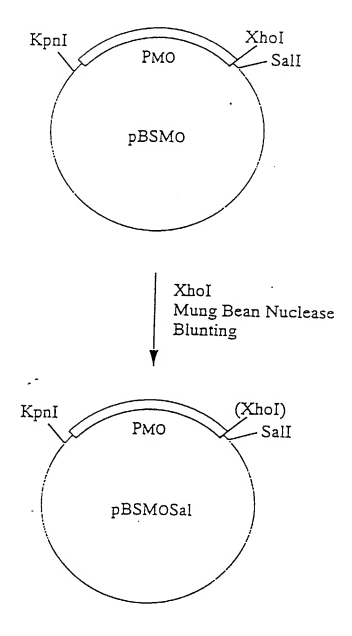


FIG.31

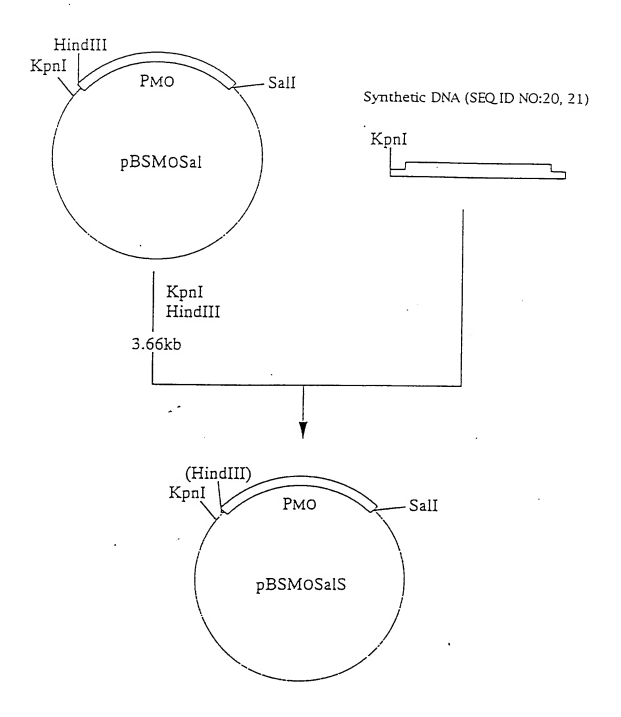


FIG.32

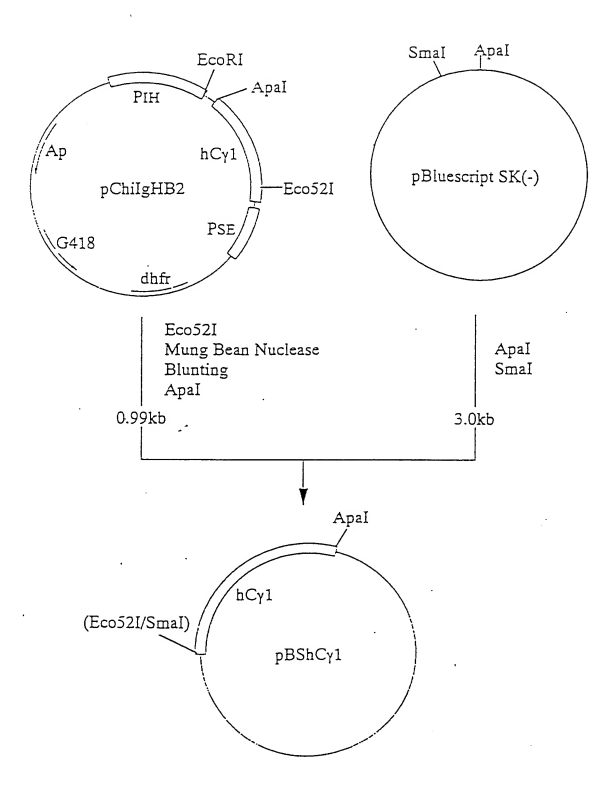
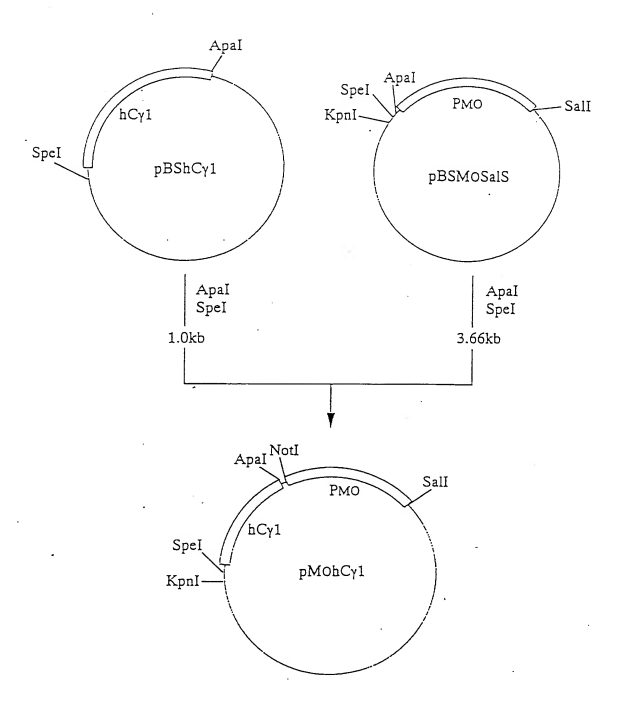


FIG.33



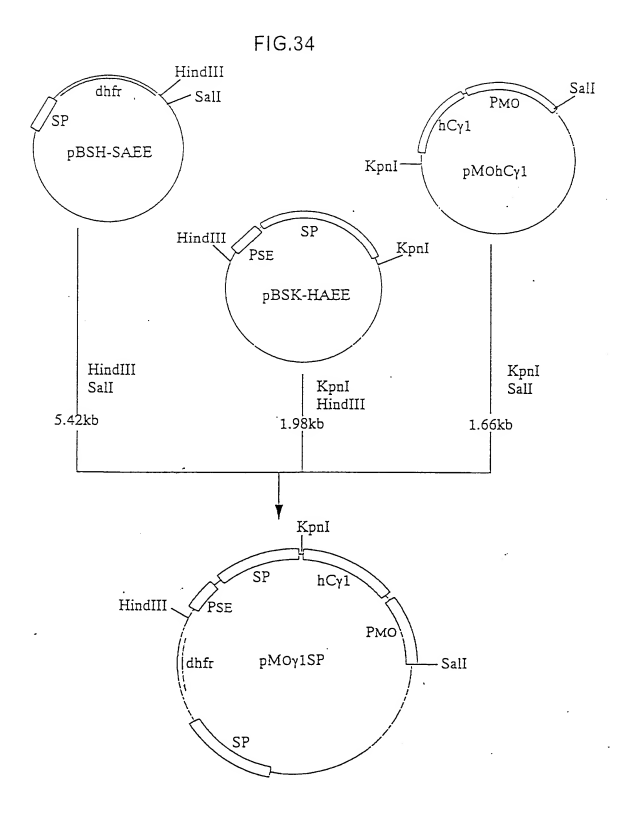


FIG.35

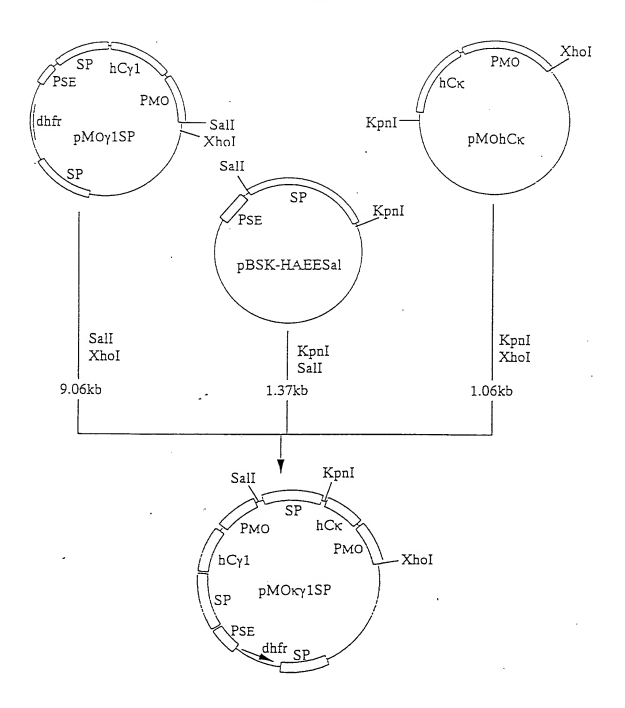


FIG.36

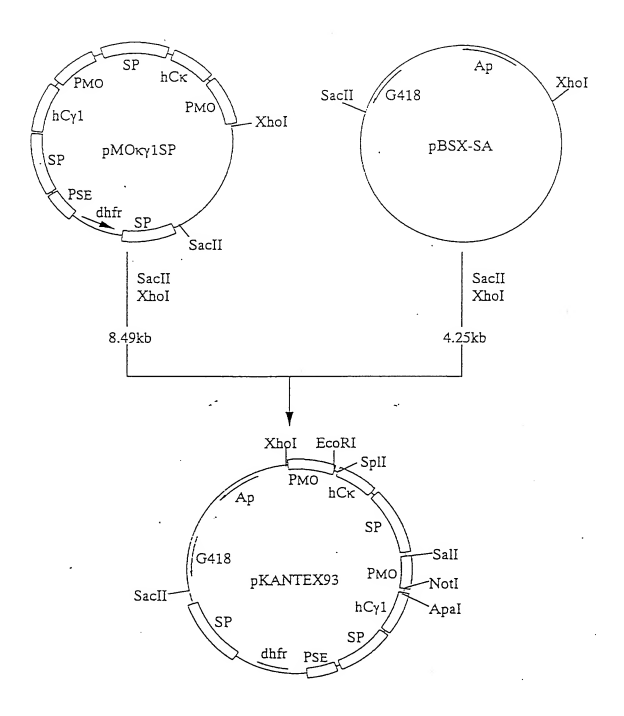


FIG.37

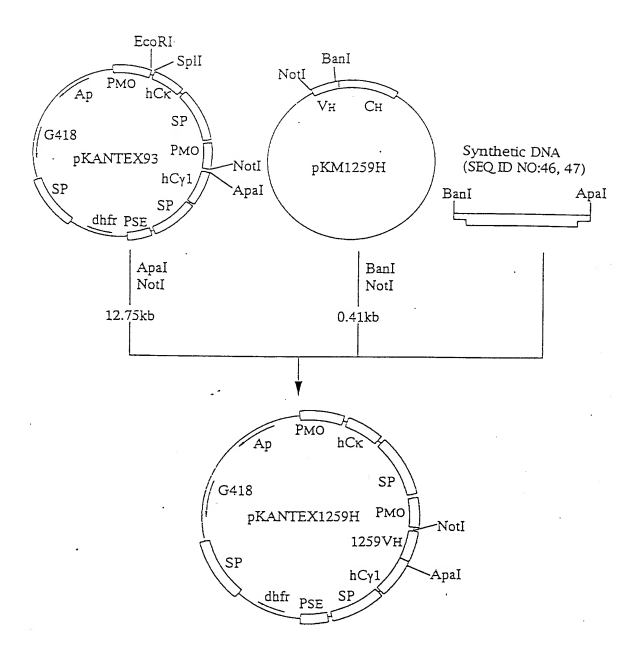


FIG.38

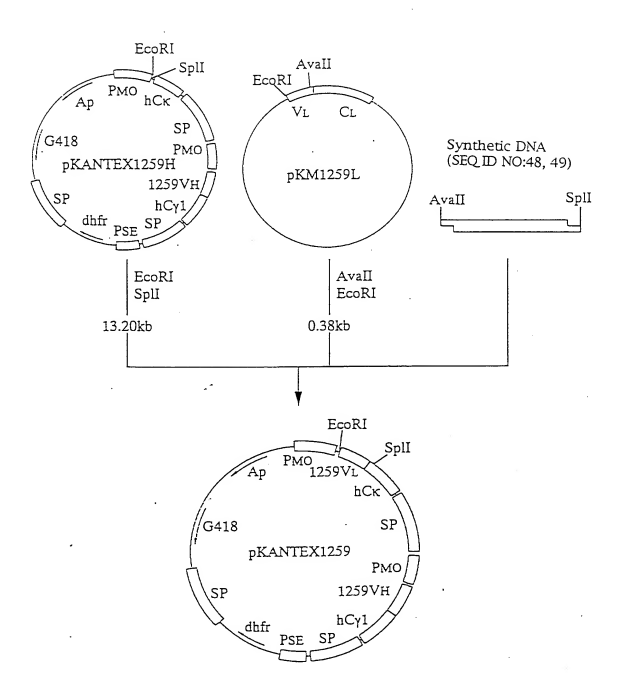


FIG.39

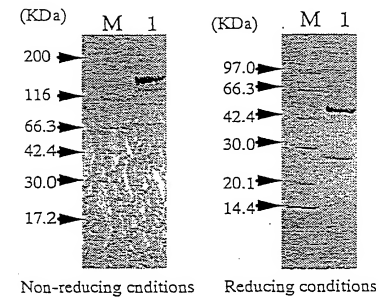


FIG.40

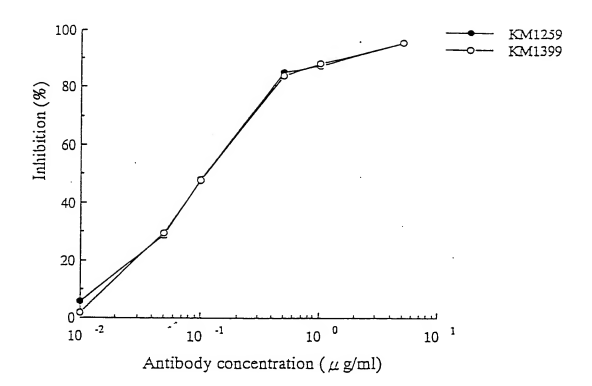


FIG.41

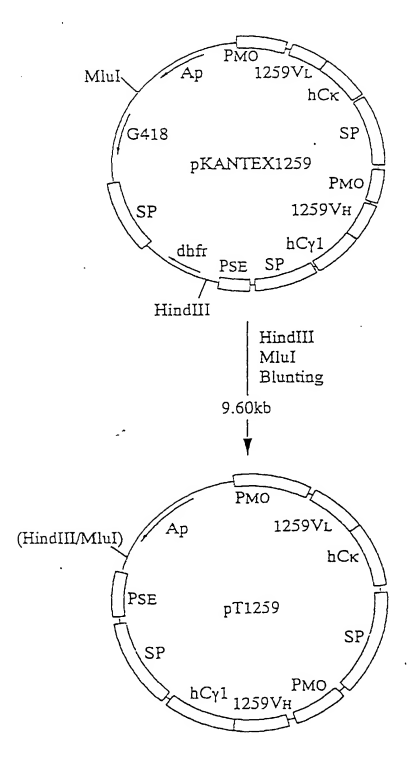


FIG.42

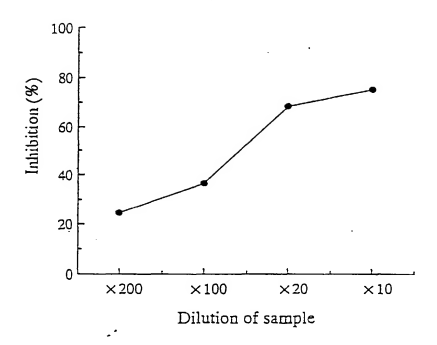


FIG.43

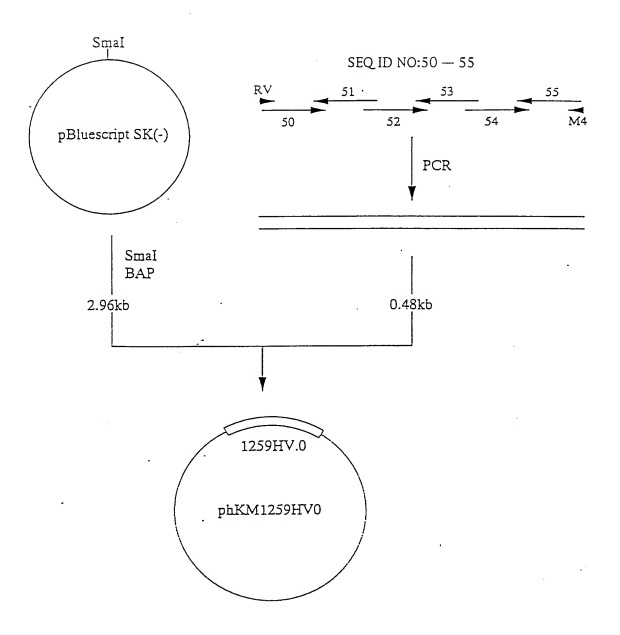


FIG.44

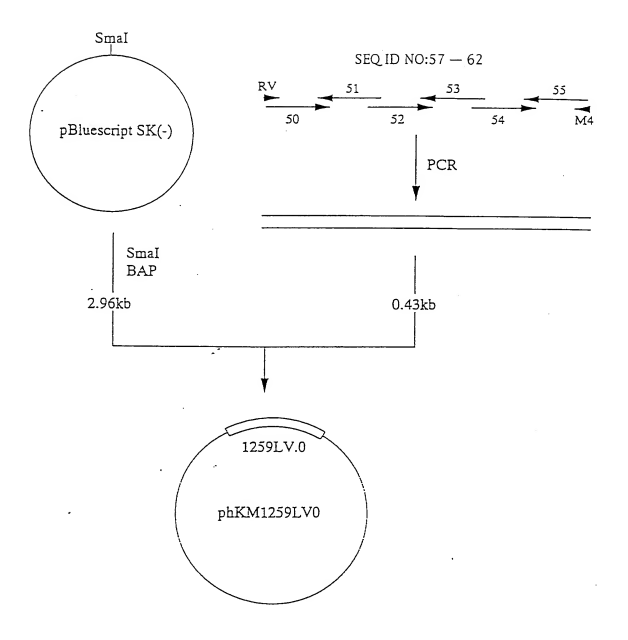


FIG.45

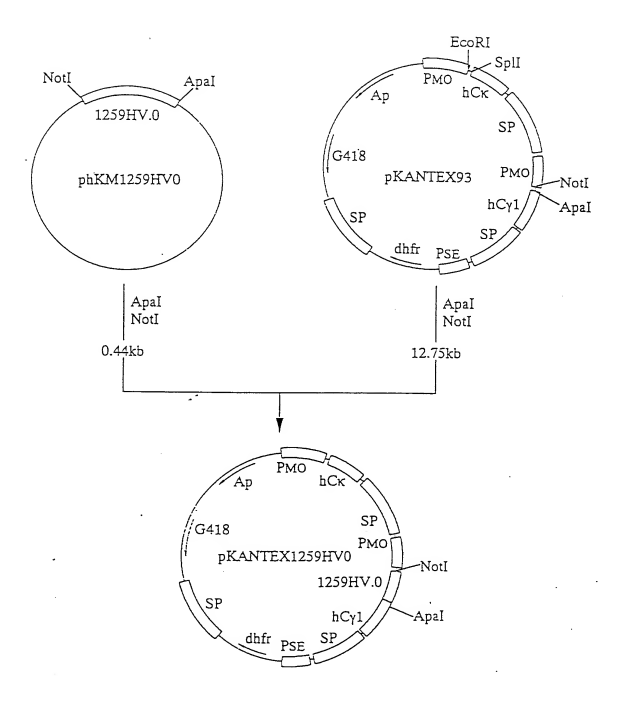


FIG.46

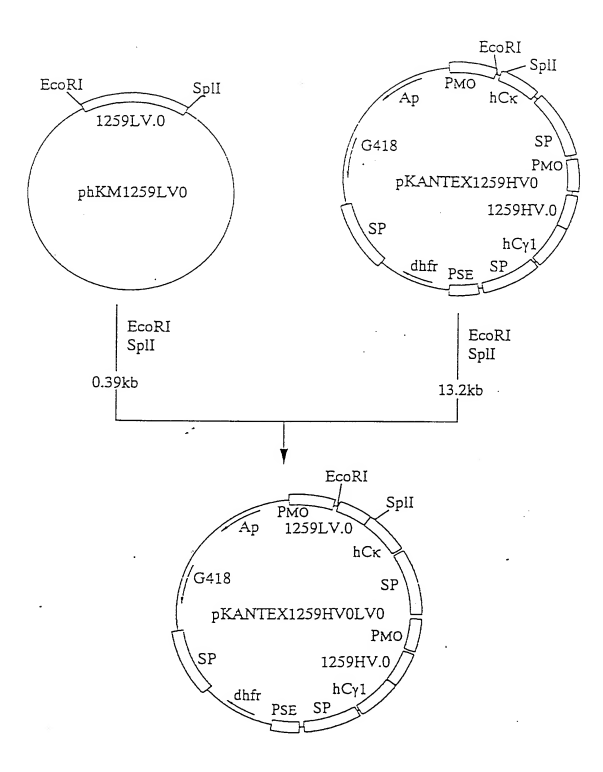
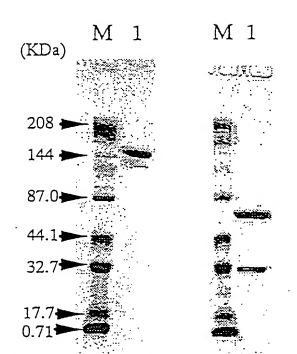


FIG.47



Non-reducing conditions Reducing conditions

FIG.48

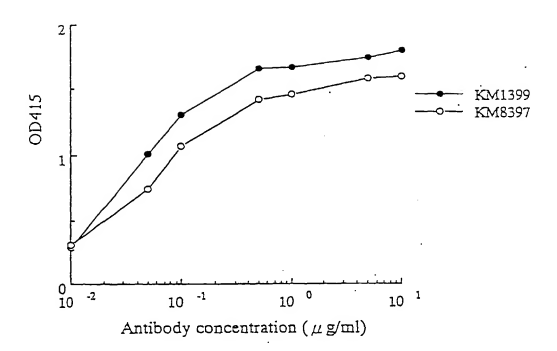


FIG.49

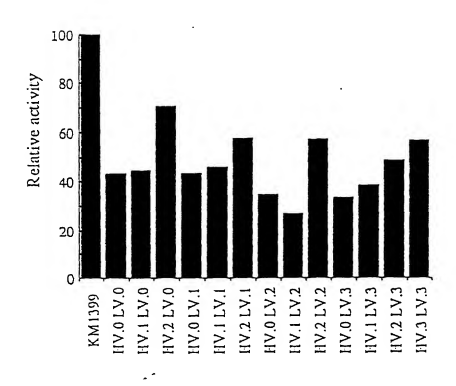
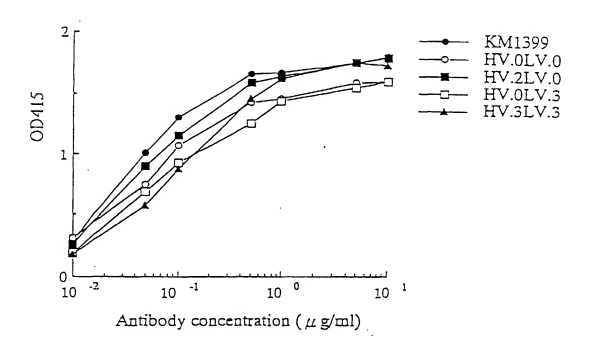


FIG.50



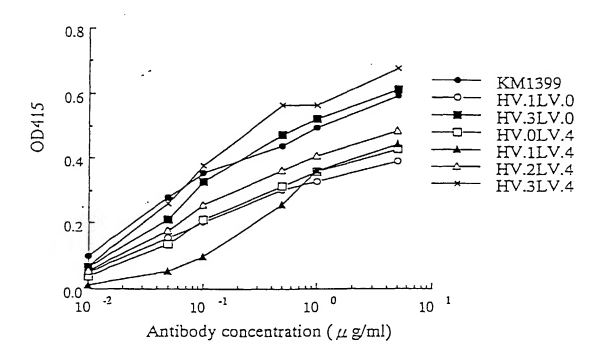


FIG.51

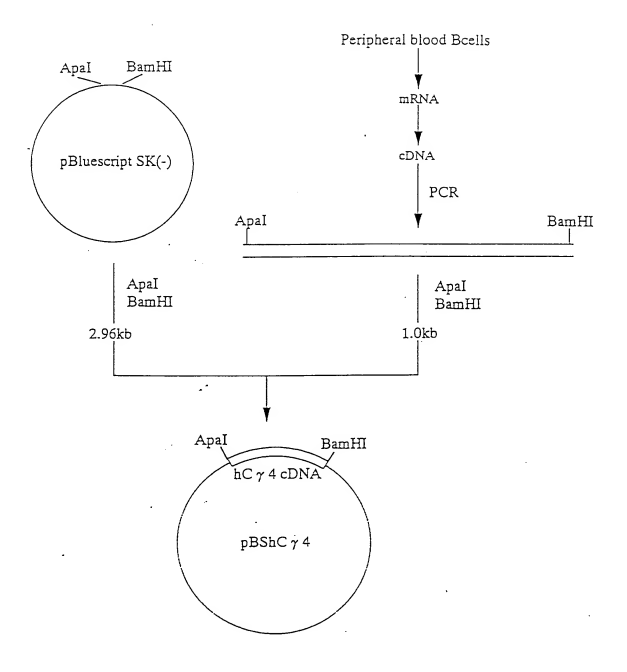


FIG.52

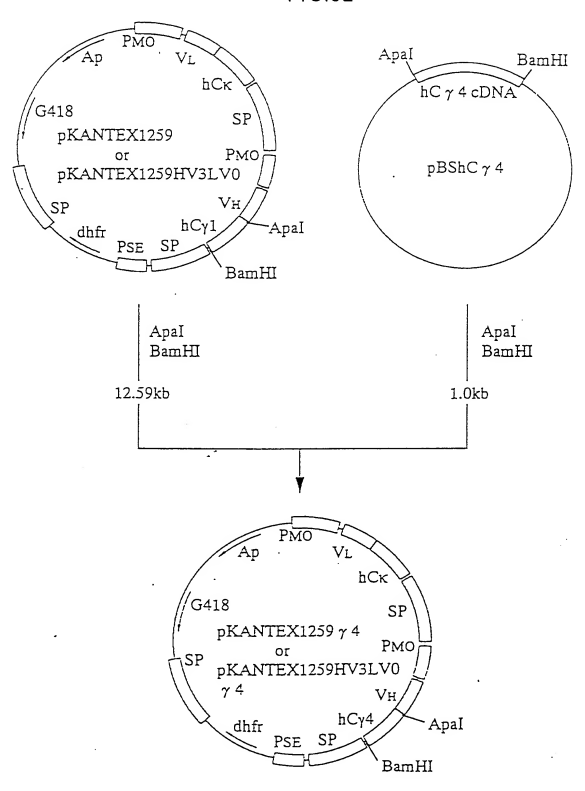


FIG.53

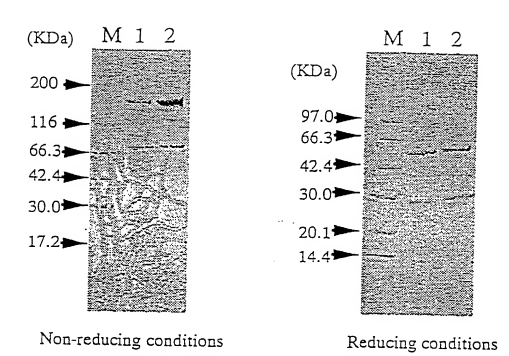


FIG.54

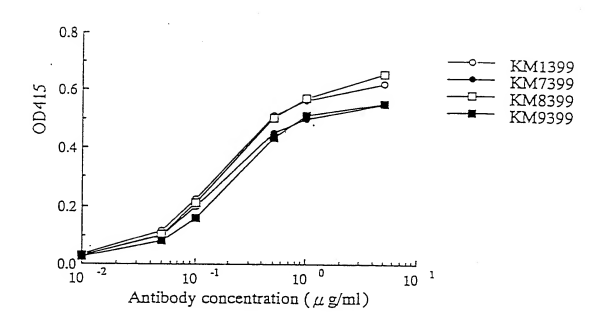


FIG.55

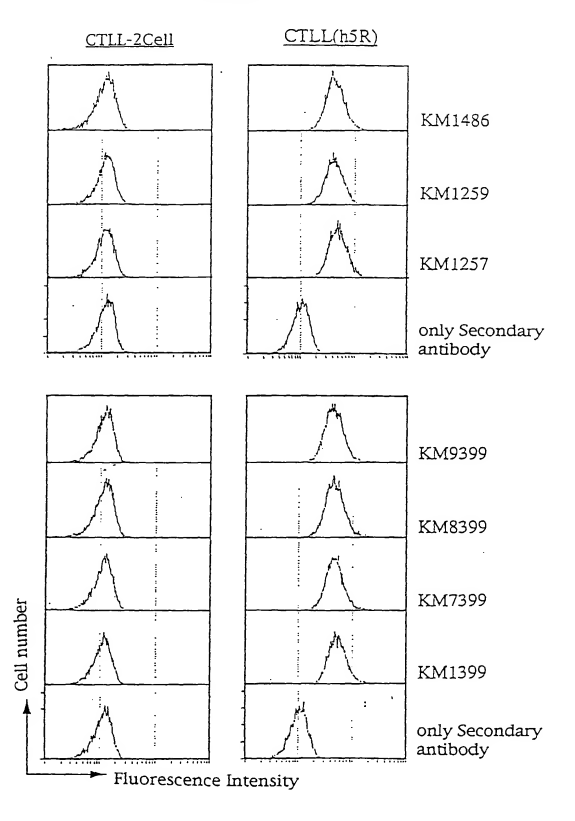


FIG.56

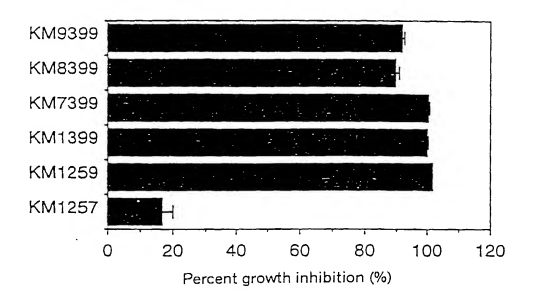
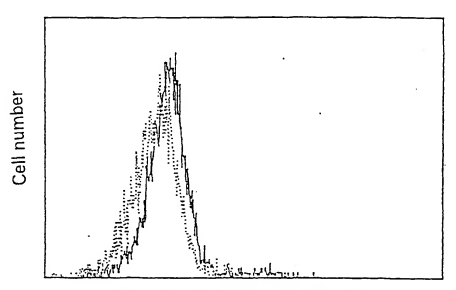


FIG.57



Relative fluorescence intensity

Dashed line: Control antibody (KM341)

Solid line : Anti-Human IL-5R α monoclonal antibody (KM1259)

FIG.58

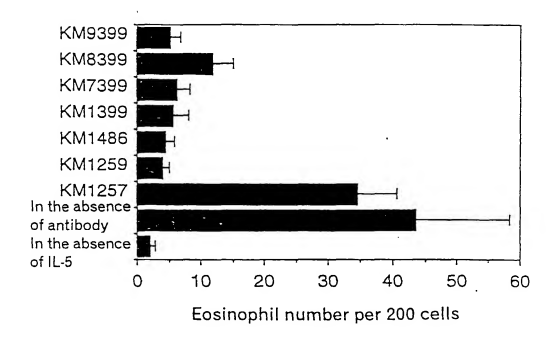


FIG.59

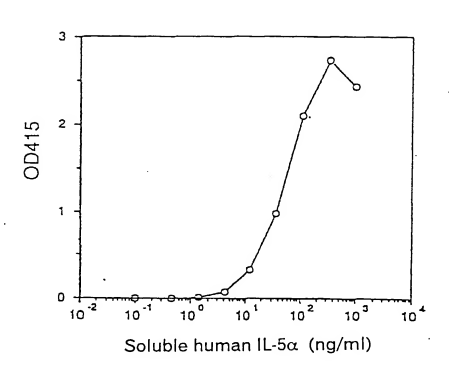
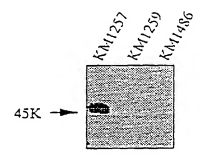


FIG.60

In the absence of 2-mercaptoethanol



In the presence of 2-mercaptoethanol

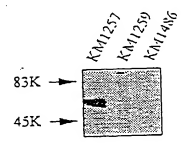
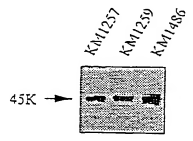


FIG.61

Immunoprecipetation of human IL-5 receptor α chain



INTERNATIONAL SEARCH REPORT

International application No.

			. PCT/J	P96/02588	
A. CLASSIFICATION OF SUBJECT MATTER Int. C1 ⁶ C12P21/08, C12N15/06, C12N15/09, C12N15/139, C12N15/63, C12N15/85, C12N5/20, C07K16/28, C07K14/715, G01N33/57, G01N33/53, A61K39/00, A61K39/395//(C12P21/08, According to International Patent Classification (IPC) or to both national classification and IPC C12R1:91)					
B. FIELDS SEARCHED					
Minimum documentation searched (classification system followed by classification symbols) Int. C1 ⁶ C12N15/02-15/90, C12P21/00, 21/02, 21/08					
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched					
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) BIOSIS PREVIEWS, WPI, WPI/L					
C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where ap	propriate, of the relev	ant passages	Relevant to claim No.	
х	Sigrid Cornelis, et al., "Detailed analysis of the IL-5-IL-5R a interaction: characterization of crucial residues on the ligand and the receptor", EMBO. J. (July 1995) Vol. 14, No. 14 p. 3395-3402			1-3, 15, 19, 20, 27, 50-54, 57, 58	
Y				4, 6, 7, 12, 16, 18, 37, 42, 47	
х	H. DeFex L., et al., "Characterization of anti IL-5 receptor monoclonal antibodies using surface plasmon resonance", FASEB J. (April 1995) Vol. 9, No. 6, p. A1502			1-3, 15, 19, 20, 27, 50-54, 57, 58	
Y				4, 6, 7, 12, 16, 18, 37, 42, 47	
Y	Satoshi Takaki, et al., "Re functional receptors for mu	constitutio	n of the men	4, 6, 7, 12, 16, 18,	
X Further documents are listed in the continuation of Box C. See patent family annex.					
Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "T" later document published after the international filing date or priori date and not in conflict with the application but cited to understant to be of particular relevance "T" later document published after the international filing date or priori date and not in conflict with the application but cited to understant to be of particular relevance.					
"L" docume	document but published on or after the international filing date ent which may throw doubts on priority claim(s) or which is o establish the publication date of another citation or other	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone			
special "O" docume means	reason (as specified) ent referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art			
	ent published prior to the international filing date but later than prity date claimed	_	"&" document member of the same patent family		
Date of the actual completion of the international search Date of mailing of the international search report					
	ember 13, 1996 (13. 12. 96)	December 25, 1996 (25. 12. 96)			
	nailing address of the ISA/	Authorized officer			
Jap.	anese Patent Office	Telephone No.			
racsimile No.		Telephone 140.			

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EP 0 811 691 A1

INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP96/02588

C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT	
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	interleukin 5", J. Exp. Med. (1993) Vol. 177, No. 6, p. 1523-1529	47-49
Y	Yoshiyuki Murata, et al., "Molecular cloning and expression of the human interleukin 5 receptor", J. Exp. Med. (1992) Vol. 175, No. 2, p. 341-351	4, 6, 7, 12 16, 18, 47-49
Y	Robert E. Bird, et al., "Single-chain antigen- binding proteins", Science (1988) Vol. 242, p. 423-426	37
Y	Keith O. Webber, et al., "Preparation and characterization of a disulfide-stabilized Fv fragment of the anti-Tac antibody: comparison with its single-chain analog", Molecular Immunology (1995) Vol. 32, No. 4, p. 249-258	42
Y	Keith O. Webber, et al., "Rapid and specific uptake of anti-Tac disulfide-stabilized Fv by interleukin-2 receptor-bearing tumors", Cancer Research (January 1995) Vol. 55, No. 2, p. 318-323	37, 42
A	M. Korenaga, et al., "The role of interleukin-5 in protective immunity to strongyloides venezuelesis infection in mice", Immunology (1991) Vol. 72, No. 4, p. 502-507	5, 8-11, 13 14, 17, 21-26, 28-3 38-41, 43-4
A	JP, 2-257891, A (Kyowa Hakko Kogyo Co., Ltd.), October 18, 1990 (18. 10. 90)(Family: none)	35, 36
	÷	

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